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#### (57) Abstract

The present invention relates to a tumor suppressor gene, termed large tumor suppressor (lats), and methods for identifying tumor suppressor genes. The method provides nucleotide sequences of lats genes, and amino acid sequences of their encoded proteins, as well as derivatives (e.g., fragments) and analogs thereof. In a specific embodiment, the lats protein is a human protein. The invention further relates to fragments (and derivatives and analogs thereof) of lats which comprise one or more domains of a lats protein. Antibodies to lats, its derivatives and analogs, are additionally provided. Methods of production of the lats proteins, derivatives and analogs, e.g., by recombinant means, are also provided. Therapeutic and diagnostic methods and pharmaceutical compositions are provided. The invention also relates to recombinant plants and animals and methods of increasing the growth of edible plants and animals. In specific examples, isolated lats genes, from Drosophila, mouse, and human, and the sequences thereof, are provided.

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### NUCLEOTIDE AND PROTEIN SEQUENCES OF LATS GENES AND METHODS BASED THEREON

#### 1. INTRODUCTION

This application is a continuation-in-part of 5 copending application Serial No. 08/411,111 filed March 27, 1995, which is incorporated by reference herein in its entirety.

The present invention relates to tumor suppressor genes, in particular to "lats" genes (large tumor suppressor)

10 and their encoded protein products, as well as derivatives and analogs thereof. Production of lats proteins, derivatives, and antibodies is also provided. The invention further relates to therapeutic compositions and methods of diagnosis and therapy.

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#### 2. BACKGROUND OF THE INVENTION

Tumorigenesis in humans is a complex process involving activation of oncogenes and inactivation of tumor suppressor genes (Bishop, 1991, Cell 64:235-248). Tumor

- 20 suppressor genes in humans have been identified through studies of genetic changes occurring in cancer cells (Ponder, 1990, Trends Genet. 6:213-218; Weinberg, 1991, Science 254:1138-1146). In Drosophila, tumor suppressor genes have been previously identified by recessive overproliferation
- 25 mutations that cause late larval and pupal lethality (Gateff, 1978, Science 200:1448-1459; Gateff and Mechler, 1989, CRC Crit. Rev. Oncogen 1:221-245; Bryant, 1993, Trends Cell Biol. 3:31-35; Török et al., 1993, Genetics 135:71-80). Mutations of interest were identified when dissection of dead larvae
- 30 and pupae revealed certain overproliferated tissues. Several genes identified in homozygous mutants have been cloned including 1(1)discs large-1(dlg; Woods and Bryant, 1991, Cell 66:451-464; Woods and Bryant, 1993, Mechanisms of Development 44:85-89), fat (Mahoney et al., 1991, Cell 67:853-868),
- 35 1(2)giant larvae (lgl. Lützelschwab et al., 1987, EMBO J. 6:1791-1797; Jacob et al., 1987, Cell 50:215-225), expanded (ex; Boedigheimer and Laughon, 1993, Development

118:1291-1301; Boedigheimer et al., 1993, Mechanisms of Development 44:83-84), hyperplastic discs (hyd; Mansfield et al., 1994, Developmental Biology 165:507-526) and the gene encoding the S6 ribosomal protein (Watson et al., 1992, Proc.

5 Natl. Acad. Sci. USA 89:11302-11306; Stewart and Denell, 1993, Mol. Cell. Biol. 13:2524-2535).

Although examining homozygous mutant animals has allowed the successful identification of overproliferation mutations that cause late larval and pupal lethality,

- 10 mutations that cause lethality at early developmental stages are unlikely to be recovered by this approach. The present invention solves this problem by providing a method for identifying tumor suppressor genes that does not exclude genes that when mutated cause lethality in early
- 15 developmental stages, and provides genes thus identified with a fundamental role in regulation of cell proliferation.

The cessation of proliferative capacity by cells in culture is termed cellular senescence. Cellular senescence is used as an experimental model for cellular aging. Normal

- 20 vertebrate cells in culture have a finite lifespan in that they undergo a characteristic maximum number of population doublings. The maximum number of population doublings that a cell can undergo inversely correlates with the age of the human donor. Cells from many human tumors are immortal cell
- 25 lines when grown in tissue culture, i.e., they exhibit infinite or continuous cell growth, suggesting that overcoming senescence is part of carcinogenesis. (For the foregoing see Hubbard and Ozer, 1995, "Senescence and immortalization of human cells," in <u>Cell Growth and</u>
- 30 Apoptosis, A Practical Approach, Ch. 12, Studzinski, G.P. (ed.), Oxford University Press Inc., New York, NY, pp. 229-248; Hubbard-Smith et al., 1992, Mol. Cell. Biol. 12:2273-2281). A comparative study of preimmortalized and immortalized human fibroblasts transformed with a defective
- 35 SV40 genome has led to the suggestion that a chromosomal region at and/or distal to 6q21 plays a role in



immortalization of cells (Hubbard-Smith et al., 1992, Mol. Cell. Biol. 12:2273-2281).

Citation of references hereinabove shall not be construed as an admission that such references are prior art to the present invention.

#### 3. SUMMARY OF THE INVENTION

The present invention relates to nucleotide sequences of lats genes (Drosophila, human, and mouse lats

10 and lats homologs of other species), and amino acid sequences of their encoded proteins, as well as derivatives (e.g., fragments) and analogs thereof. Nucleic acids hybridizable to or complementary to the foregoing nucleotide sequences are also provided. In a specific embodiment, the lats protein is a human protein.

The invention also relates to a method of identifying tumor suppressor genes that does not exclude from identification genes that cause lethality at early developmental stages, thus overcoming the limitations of prior art methods. The method thus allows the identification of genes that regulate cell proliferation and that act at early developmental stages. The genes which thus can be identified play a fundamental role in regulation of cell proliferation such that their dysfunction (e.g., by lack of expression or mutation) leads to overproliferation and cancer.

Lats is a gene provided by the present invention, identified by the method of the invention, that acts to inhibit cell proliferation, and that plays a crucial role throughout development.

The invention also relates to lats derivatives and analogs of the invention which are functionally active, i.e., they are capable of displaying one or more known functional activities associated with a full-length (wild-type) lats protein. Such functional activities include but are not limited to kinase activity, antigenicity [ability to bind (or compete with lats for binding) to an anti-lats antibody],

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immunogenicity (ability to generate antibody which binds to lats), and ability to bind (or compete with lats for binding) to a receptor/ligand for lats (e.g., a SH3 domain-containing protein).

The invention further relates to fragments (and derivatives and analogs thereof) of lats which comprise one or more domains of a lats protein.

Antibodies to lats, and lats derivatives and analogs, are additionally provided.

Methods of production of the lats proteins, derivatives and analogs, e.g., by recombinant means, are also provided.

The present invention also relates to therapeutic and diagnostic methods and compositions based on lats

15 proteins and nucleic acids. Therapeutic compounds of the invention include but are not limited to lats proteins and analogs and derivatives (including fragments) thereof; antibodies thereto; nucleic acids encoding the lats proteins, analogs, or derivatives; and lats antisense nucleic acids.

- The invention provides for treatment of disorders of overproliferation (e.g., cancer and hyperproliferative disorders) by administering compounds that promote lats activity (e.g., lats, an agonist of lats; nucleic acids that encode lats).
- The invention also provides methods of treatment of disorders involving deficient cell proliferation (growth) or in which cell proliferation is otherwise desired (e.g., degenerative disorders, growth deficiencies, lesions, physical trauma) by administering compounds that antagonize, (inhibit) lats function (e.g., antibodies, antisense nucleic acids).

In a specific embodiment, lats function is antagonized in order to inhibit cellular senescence, in vivo or in vitro.

Antagonizing lats function can also be done to grow larger animals and plants, e.g., those used as food or material sources.



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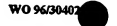
Animal models, diagnostic methods and screening methods for predisposition to disorders, and methods to identify lats agonists and antagonists, are also provided by the invention.

3.1. <u>DEFINITIONS</u>

As used herein, underscoring or italicizing the name of a gene shall indicate the gene, in contrast to its encoded protein product which is indicated by the name of the 10 gene in the absence of any underscoring or italicizing. For example, "lats" shall mean the lats gene, whereas "lats" shall indicate the protein product of the lats gene.

#### 4. DESCRIPTION OF THE FIGURES

- Pigure 1. Identifying overproliferation mutations in mosaic flies. (A) Although animals that are homozygous for a lethal mutation could die at an early developmental stage, mosaic flies carrying clones of cells that are homozygous for the same mutation could live. One can identify potential
- 20 tumor suppressors by generating and examining clones of overproliferated mutant cells in mosaic animals. The genetic constitution of these mosaic flies is similar to the mosaicism of the tumor patients. (B) Genetic scheme. The P-element insertions carrying the FLP recombinase (hsFLP;
- 25 Golic and Lindquist, 1989, Cell 59:499-509), its target site, FRT (solid arrows, Xu and Rubin, 1993, Development 117:1223-1237), the yellow\* and mini-white\* marker genes (y\* and mini-w\*, open arrows) are indicated. Mutagenized males were crossed to females to produce heterozygous embryos.
- 30 Clones of cells homozygous for the induced mutations were generated in developing first-instar larvae by mitotic recombination at the FRT sites induced with the FLP recombinase. Mosaic adults were examined for overproliferated mutant patches (w, y). Individuals
- carrying clones of interest were then mated to recover the mutations of interest in the next generation (Xu and Rubin,



1993, Development 117:1223-1237; Xu and Harrison, 1994; Methods in Cell Biology 44:655-682). Clones of ommatidia derived from fast proliferating mutant cells were identified since they were larger than their darkly pigmented wt (wild-type) twin-spot clones (mini-w\*/mini-w\*).

Figure 2. Mutant phenotypes. (A) A clone of unpatterned, overproliferated lats mutant cells in the eye. (B) Induced at the same stage, the 93B mutant cells formed a less overproliferated clone. (C) A third instar lats 26-1 larva 10 (right) was much larger than a wt sibling (left; at 18°C). (D) Wing discs from the larva in (C) (wt, top; lats 261, bottom). (E) Dissected central nervous systems (wt, top; latsell, bottom). (F) A SEM (scanning electron microscope) view of a lats clone near the eye. (G) A closer view of a 15 region in (F) showing the irregularity of the sizes and shapes of the mutant cells. (H) A plastic section of a mutant clone similar to the one in (F). Cells seem to be "budding" out of the surface to form new proliferating lobes (arrows). (I) A lats clone on the back. The boxed area is 20 shown in (J). The bristles in the mutant clone are short, bent and often split (arrows). (K) A closer view of the hairs in a lats clone on the body showing enlarged bases and bent tips. (L) A section of a lats clone on the back showing extra cuticle deposits (arrows). All the mutant clones were 25 induced with lats" unless stated differently.

gene. The genomic restriction map of the lats region is aligned with the lats 5.7 kb transcript unit. The direction of transcription is indicated with large arrows. The sizes 30 of the lats introns are as follows: intron 1 (5.0 kb), intron 2 (5.8 kb), intron 3 (68 bp), intron 4 (63 bp), intron 5 (64 bp), intron 6 (61 bp), intron 7 (62 bp). The genomic DNA from +7.5 (BglII) to -4.2 (EcoRI) was used to screen a total imaginal disc cDNA library, which isolated three groups of cDNAs: lats, T1, T2. The introns in the T2 transcript are not labeled. Only parts of the zfh-1 (Fortini et al., 1991, Mechan. Dev. 34:113-122) and T1 transcripts are

Figure 3. Organization of the Drosophila lats



indicated. The locations of the P-element insertion ( $lats^{Pl}$ ), the deletions in the five excision alleles ( $lats^{e7-2, e78, e100, e119, e148}$ ) and in  $lats^{el}$ ,  $lats^{el}$  are indicated at the bottom. The slash indicates a gap in the genomic map. Restriction sites:

5 EcoRI (small open arrow), BglII (open box) and BamHI (open circle). The BglII site at the -0.5 position of the CLT-52 clone is not present in other genomic DNA. A scale is labeled under the restriction map.

Figure 4. RNA blot analysis of the Drosophila lats

10 mRNA. Five μg of poly(A) \* RNA isolated from various developmental stages was separated on a 1% agarose gel, and hybridized with <sup>32</sup>P-labeled 5 \* end 1 kb probe from the Drosophila lats cDNA. E0-2 hrs, E2-4 hrs, E4-6 hrs, E6-8 hrs, E8-16 hrs and E16-24 hrs indicate the age of the embryos

- 15 in hours. RNA from first, second and third instar larvae is denoted by L1, L2, and L3, respectively. The numbers and arrows on the right correspond to the size and location of the RNA standards. A 5.7 kb RNA was found in all the developmental stages, whereas a 4.7 kb RNA was predominantly
- 20 present in 0 to 4 hour old embryos. The blot was also hybridized with DNA from the ribosomal protein gene, RNA1.

Figure 5. Composite cDNA sequence of the Drosophila lats gene. The entire cDNA sequence (SEQ ID NO:1) corresponding to the 5.7 kb lats RNA is shown. This

- nucleotide sequence is a composite of two cDNA clones (nucleotide 1-191 from cDNA 9 and the rest from cDNA A2). The sequence of the corresponding genomic DNA has been determined and is identical to the cDNA sequence except where indicated (above the cDNA sequence). The predicted amino
- 30 acid sequence (SEQ ID NO:2) is shown below the cDNA sequence. The opa repeat is indicated by the heavy bar. The location of the putative SH3 binding site and the RERDQ peptides are designated by dashed lines. The two sites that match the polyadenylation signal consensus sequence are underlined.
- 35 The second site is located at 12 bp away from the 3' end of the cDNA. The locations of the introns are indicated by vertical arrows. The underlined 141 bp sequence at the 3'

end of the lats transcript is identical to the 5' end untranslated sequence of the class I transcript of the Drosophila phospholipase C gene, plc-21. The location of the 446 bp deletion in the lats<sup>al</sup> allele is also indicated.

- Figure 6. Schematic of the Drosophila lats predicted protein (SEQ ID NO:2) and the related proteins (A) and sequence comparison of the proteins homologous to lats (B). In Fig. 6A, solid, hatched, open and shaded boxes denote putative SH3 binding site, opa repeat, RERDQ peptide
- 10 and kinase domain in the lats protein, respectively. The Dbf20, Dbf2 and COT-1 proteins are illustrated at the bottom. The regions that are homologous to lats are indicated by shaded boxes. The degrees of sequence similarity (percentage of identical sequences inside parentheses; percentage of
- 15 identical or conservative substitutions outside parentheses) between lats and the three related proteins are indicated above the corresponding regions of these proteins. In Fig. 6B, the carboxy-terminal half of lats is compared to the six most related proteins that are revealed by blastp (a software
- 20 program that searches for protein sequence homologies) search
  as of Sept. 1, 1994. Neurospora cot-1 (SEQ ID NO:11);
  tobacco PKTL7 (SEQ ID NO:12); common ice plant protein kinase
  (SEQ ID NO:13); spinach protein kinase (SEQ ID NO:14); yeast
  Dbf-20 (SEQ ID NO:15); yeast Dbf2 (SEQ ID NO:16). Amino acid
- 25 residues identical to lats are highlighted. Numbers at the beginning of every sequence refer to the position of that amino acid within the total protein sequence. The boundary of the kinase domain is defined according to Hanks et al. (1988, Science 241:42-52). The location of a region of about
- 30 40 amino acid residues that is not conserved among the proteins is indicated by the heavy bar above the sequence. The sequence of PKTL7 from tobacco, Nicotiana tabacum, was submitted to Genbank by Huang, Y. (X71057). Both the sequence of the protein kinase from spinach, Spinacia oleracea, and
- 35 the sequence of the protein kinase from common ice plant,

  Mesembryanthemum crystallinum, were submitted to Genbank by



Baur, B., Winter, K., Fischer, K. and Dietz, K. (Z30329 and Z30330).

Figure 7. cDNA sequence (SEQ ID NO:5) and deduced protein sequence (SEQ ID NO:6) of a mouse lats homolog, 5 m-lats.

Figure 8. cDNA sequence (SEQ ID NO:7) and deduced protein sequence (SEQ ID NO:8) of a mouse lats homolog, m-lats2.

Figure 9. cDNA sequence (SEQ ID NO:3) and deduced 10 protein sequence (SEQ ID NO:4) of a human lats homolog, h-lats.

Figure 10. Schematic diagram of plasmid pBS(KS)-h-lats, containing the full length coding sequence of the h-lats cDNA.

- sequence (SEQ ID NO:4) (upper case letters) with the m-lats protein sequence (SEQ ID NO:6) (lower case letters). A dot indicates amino acid identity; a dash indicates a deletion relative to the sequence on the line above. The amino-
- 20 terminal portion of the m-lats protein is not shown due to the missing 5' end of the m-lats cDNA coding region.

Figure 12. Alignment of the h-lats protein sequence (SEQ ID NO:4) (upper case letters) with the m-lats2 protein sequence (SEQ ID NO:8) (lower case letters). A dot

25 indicates amino acid identity; a dash indicates a deletion relative to the sequence on the line above. The aminoterminal portion of the m-lats2 protein is not shown due to the missing 5' end of the m-lats2 cDNA coding region.

Figure 13. Alignment of the h-lats protein

30 sequence (SEQ ID NO:4) (upper case letters) with the

Drosophila lats protein sequence (SEQ ID NO:2) (lower case
letters). A dot indicates amino acid identity; a dash
indicates a deletion relative to the sequence on the line
above. Insertions in the Drosophila sequence relative to the

35 human sequence are indicated below the sequence line.

Conserved domains are indicated. LSD2 = lats split domain 2;

LSD2a = LSD2 anterior portion; LSD2p = LSD2 posterior

portion. The putative SH3-binding domain and the kinase domain are shown. LSD1 = lats split domain 1; LSD1a = LSD1 anterior portion; LSD1p = LSD1 posterior portion. LFD = lats flanking domain. LCD1 = lats C-terminal domain 1; LCD2 = lats C-terminal domain 2; LCD3 = lats C-terminal domain 3.

Figure 14. Schematic diagram of plasmid pCaSpeR-hs-h-lats, an expression vector containing the full length coding sequence of the h-lats cDNA.

Pigure 15. Northern blot analysis of h-lats

10 expression in normal human tissues. A  $^{32}$ P-labeled BamHI fragment of h-lats was used as a probe for hybridization to polyA+ RNA from the normal human fetal and adult tissues indicated for each lane. The positions of standard molecular weight markers are shown at right. The positions of the

15 h-lats RNA and of  $\beta$ -actin RNA (used as a standard) are shown.

#### 5. DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to nucleotide sequences of lats genes, and amino acid sequences of their 20 encoded proteins. The invention further relates to fragments and other derivatives, and analogs, of lats proteins.

Nucleic acids encoding such fragments or derivatives are also within the scope of the invention. The invention provides lats genes and their encoded proteins of many different

- 25 species. The lats genes of the invention include Drosophila, human, and mouse lats and related genes (homologs) in other species. In specific embodiments, the lats genes and proteins are from vertebrates, or more particularly, mammals. In a preferred embodiment of the invention, the lats genes
- 30 and proteins are of human origin. Production of the foregoing proteins and derivatives, e.g., by recombinant methods, is provided.

The invention also relates to a method of identifying tumor suppressor genes that does not exclude from 35 identification genes that cause lethality at early developmental stages, thus overcoming the limitations of prior art methods. The method thus allows the identification



of genes that regulate cell proliferation and that act at early developmental stages. The genes which thus can be identified play a fundamental role in regulation of cell proliferation such that their dysfunction (e.g., due to lack of expression or mutation) leads to overproliferation and cancer.

Lats is a gene provided by the present invention, identified by the method of the invention, that acts to inhibit cell proliferation, and that plays a crucial role throughout development.

The invention also relates to lats derivatives and analogs of the invention which are functionally active, i.e., they are capable of displaying one or more known functional activities associated with a full-length (wild-type) lats protein. Such functional activities include but are not limited to kinase activity, antigenicity (ability to bind (or compete with lats for binding) to an anti-lats antibody), immunogenicity (ability to generate antibody which binds to lats), ability to bind (or compete with lats for binding) to an SH3-domain-containing protein or other ligand, ability to

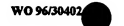
The invention further relates to fragments (and derivatives and analogs thereof) of lats which comprise one or more domains of the lats protein.

The present invention also relates to therapeutic

Antibodies to lats, its derivatives and analogs, are additionally provided.

inhibit cell proliferation, tumor inhibition, etc.

and diagnostic methods and compositions based on lats proteins and nucleic acids and anti-lats antibodies. The invention provides for treatment of disorders of overproliferation (e.g., cancer and hyperproliferative disorders) by administering compounds that promote lats activity (e.g., lats proteins and functionally active analogs and derivatives (including fragments) thereof; nucleic acids encoding the lats proteins, analogs, or derivatives, agonists of lats).



The invention also provides methods of treatment of disorders involving deficient cell proliferation or in which cell proliferation (growth) is otherwise desirable (e.g., growth deficiencies, degenerative disorders, lesions,

5 physical trauma) by administering compounds that antagonize, or inhibit, lats function (e.g., antibodies, lats antisense nucleic acids, lats derivatives that are dominant-negative protein kinases).

In a specific embodiment, lats function is

10 antagonized in order to inhibit cellular senescence, in vivo
or in vitro.

Inhibition of lats function can also be done to grow larger farm animals and plants.

Animal models, diagnostic methods and screening

15 methods for predisposition to disorders are also provided by
the invention.

The invention is illustrated by way of examples infra which disclose, inter alia, the cloning and characterization of D. melanogaster lats (Section 6); the cloning and characterization of mouse and human lats homologs (Section 7); the sequence and domain conservation among the lats homologs (Section 8); the functional interchangeability of the human and Drosophila lats homologs (Section 9); and

For clarity of disclosure, and not by way of limitation, the detailed description of the invention is divided into the subsections which follow.

the differentially decreased expression of human lats in

25 human tumor cell lines (Section 10).

30

#### 5.1. <u>ISOLATION OF THE LATS GENES</u>

The invention relates to the nucleotide sequences of lats nucleic acids. In specific embodiments, lats nucleic acids comprise the cDNA sequences of SEQ ID NO:1, 3, 5, or 7, or the coding regions thereof, or nucleotide sequences acids encoding a lats protein (e.g., a protein having the sequence of SEQ ID NO:2, 4, 6, or 8). The invention provides purified nucleic acids consisting of at least 8 nucleotides (i.e., a



hybridizable portion) of a *lats* sequence; in other embodiments, the nucleic acids consist of at least 25 (continuous) nucleotides, 50 nucleotides, 100 nucleotides, 150 nucleotides, or 200 nucleotides of a *lats* sequence, or a

- 5 full-length lats coding sequence. In another embodiment, the nucleic acids are smaller than 35, 200 or 500 nucleotides in length. Nucleic acids can be single or double stranded. The invention also relates to nucleic acids hybridizable to or complementary to the foregoing sequences. In specific
- 10 aspects, nucleic acids are provided which comprise a sequence complementary to at least 10, 25, 50, 100, or 200 nucleotides or the entire coding region of a lats gene. In a specific embodiment, a nucleic acid which is hybridizable to a lats nucleic acid (e.g., having sequence SEQ ID NO:3 or 7), or to
- 15 a nucleic acid encoding a lats derivative, under conditions of low stringency is provided. By way of example and not limitation, procedures using such conditions of low stringency are as follows (see also Shilo and Weinberg, 1981, Proc. Natl. Acad. Sci. USA 78:6789-6792): Filters containing
- 20 DNA are pretreated for 6 h at 40°C in a solution containing 35% formamide, 5X SSC, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.1% PVP, 0.1% Ficoll, 1% BSA, and 500 μg/ml denatured salmon sperm DNA. Hybridizations are carried out in the same solution with the following modifications: 0.02% PVP, 0.02%
- 25 Ficoll, 0.2% BSA, 100 μg/ml salmon sperm DNA, 10% (wt/vol) dextran sulfate, and 5-20 X 106 cpm <sup>32</sup>P-labeled probe is used. Filters are incubated in hybridization mixture for 18-20 h at 40°C, and then washed for 1.5 h at 55°C in a solution containing 2X SSC, 25 mM Tris-HCl (pH 7.4), 5 mM EDTA, and
- 30 0.1% SDS. The wash solution is replaced with fresh solution and incubated an additional 1.5 h at 60°C. Filters are blotted dry and exposed for autoradiography. If necessary, filters are washed for a third time at 65-68°C and reexposed to film. Other conditions of low stringency which may be
- 35 used are well known in the art (e.g., as employed for crossspecies hybridizations).

In another specific embodiment, a nucleic acid which is hybridizable to a lats nucleic acid under conditions of high stringency is provided. By way of example and not limitation, procedures using such conditions of high

- 5 stringency are as follows: Prehybridization of filters containing DNA is carried out for 8 h to overnight at 65°C in buffer composed of 6X SSC, 50 mM Tris-HCl (pH 7.5), 1 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.02% BSA, and 500 μg/ml denatured salmon sperm DNA. Filters are hybridized for 48 h
- 10 at 65°C in prehybridization mixture containing 100 μg/ml denatured salmon sperm DNA and 5-20 X 10° cpm of <sup>32</sup>P-labeled probe. Washing of filters is done at 37°C for 1 h in a solution containing 2X SSC, 0.01% PVP, 0.01% Ficoll, and 0.01% BSA. This is followed by a wash in 0.1X SSC at 50°C
- 15 for 45 min before autoradiography. Other conditions of high stringency which may be used are well known in the art.

In another specific embodiment, a nucleic acid, which is hybridizable to a *lats* nucleic acid under conditions of moderate stringency is provided (see, e.g., Section 7.2).

- Nucleic acids encoding derivatives and analogs of lats proteins (see Sections 5.6 and 5.6.1), and lats antisense nucleic acids (see Section 5.8.2.2.1) are additionally provided. As is readily apparent, as used herein, a "nucleic acid encoding a fragment or portion of a
- 25 lats protein" shall be construed as referring to a nucleic acid encoding only the recited fragment or portion of the lats protein and not the other contiguous portions of the lats protein as a continuous sequence.

Fragments of lats nucleic acids comprising regions

30 conserved between (with homology to) other lats nucleic
acids, of the same or different species, are also provided.

Nucleic acids encoding one or more lats domains are provided.

Specific embodiments for the cloning of a lats gene, presented as a particular example but not by way of 35 limitation, follows:

For expression cloning (a technique commonly known in the art), an expression library is constructed by methods



known in the art. For example, mRNA (e.g., human) is
isolated, cDNA is made and ligated into an expression vector
(e.g., a bacteriophage derivative) such that it is capable of
being expressed by the host cell into which it is then
introduced. Various screening assays can then be used to
select for the expressed lats product. In one embodiment,
anti-lats antibodies can be used for selection.

In another embodiment, polymerase chain reaction (PCR) is used to amplify the desired sequence in a genomic or 10 cDNA library, prior to selection. Oligonucleotide primers representing known lats sequences can be used as primers in PCR. In a preferred aspect, the oligonucleotide primers represent at least part of the lats conserved segments of strong homology between lats of different species (e.g.,

- 15 LCD1, LCD2, kinase domain, LFD, SH3 binding domain, LSD1, and LSD2 domains; see, e.g., Section 8 infra.) The synthetic oligonucleotides may be utilized as primers to amplify by PCR sequences from a source (RNA or DNA), preferably a cDNA library, of potential interest. PCR can be carried out,
- 20 e.g., by use of a Perkin-Elmer Cetus thermal cycler and Taq polymerase (Gene Amp"). The DNA being amplified can include mRNA or cDNA or genomic DNA from any eukaryotic species. One can choose to synthesize several different degenerate primers, for use in the PCR reactions. It is also possible
- 25 to vary the stringency of hybridization conditions used in priming the PCR reactions, to allow for greater or lesser degrees of nucleotide sequence similarity between the known lats nucleotide sequence and the nucleic acid homolog being isolated. For cross species hybridization, low stringency
- 30 conditions are preferred. For same species hybridization, moderately stringent conditions are preferred. After successful amplification of a segment of a lats homolog, that segment may be molecularly cloned and sequenced, and utilized as a probe to isolate a complete cDNA or genomic clone.
- 35 This, in turn, will permit the determination of the gene's complete nucleotide sequence, the analysis of its expression, and the production of its protein product for functional



analysis, as described infra. In this fashion, additional genes encoding lats proteins and lats analogs may be identified.

The above-methods are not meant to limit the

5 following general description of methods by which clones of

lats may be obtained.

Any eukaryotic cell potentially can serve as the nucleic acid source for the molecular cloning of the lats gene. The nucleic acid sequences encoding lats can be

- 10 isolated from vertebrate, mammalian, human, porcine, bovine, feline, avian, equine, canine, as well as additional primate sources, insects, plants, etc. The DNA may be obtained by standard procedures known in the art from cloned DNA (e.g., a DNA "library"), by chemical synthesis, by cDNA cloning, or by
- 15 the cloning of genomic DNA, or fragments thereof, purified from the desired cell. (See, for example, Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, 2d Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach,
- 20 MRL Press, Ltd., Oxford, U.K. Vol. I, II.) Clones derived from genomic DNA may contain regulatory and intron DNA regions in addition to coding regions; clones derived from cDNA will contain only exon sequences. Whatever the source, the gene should be molecularly cloned into a suitable vector 25 for propagation of the gene.

In the molecular cloning of the gene from genomic DNA, DNA fragments are generated, some of which will encode the desired gene. The DNA may be cleaved at specific sites using various restriction enzymes. Alternatively, one may use DNAse in the presence of manganese to fragment the DNA,

or the DNA can be physically sheared, as for example, by sonication. The linear DNA fragments can then be separated according to size by standard techniques, including but not limited to, agarose and polyacrylamide gel electrophoresis

35 and column chromatography.

Once the DNA fragments are generated, identification of the specific DNA fragment containing the



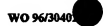
desired gene may be accomplished in a number of ways. For example, if an amount of a portion of a lats (of any species) gene or its specific RNA, or a fragment thereof (see Section 5.6), is available and can be purified and labeled, the

- 5 generated DNA fragments may be screened by nucleic acid hybridization to the labeled probe (Benton, W. and Davis, R., 1977, Science 196:180; Grunstein, M. And Hogness, D., 1975, Proc. Natl. Acad. Sci. U.S.A. 72:3961). Such a procedure is presented by way of example in Section 7 infra. Those DNA
- 10 fragments with substantial homology to the probe will hybridize. It is also possible to identify the appropriate fragment by restriction enzyme digestion(s) and comparison of fragment sizes with those expected according to a known restriction map if such is available. Further selection can
- Alternatively, the presence of the gene may be detected by assays based on the physical, chemical, or immunological properties of its expressed product. For example, cDNA clones, or DNA clones which hybrid-select the proper mRNAs,
- 20 can be selected which produce a protein that, e.g., has similar or identical electrophoretic migration, isolectric focusing behavior, proteolytic digestion maps, kinase activity, inhibition of cell proliferation activity, substrate binding activity, or antigenic properties as known
- 25 for lats. If an antibody to lats is available, the lats protein may be identified by binding of labeled antibody to the putatively lats synthesizing clones, in an ELISA (enzymelinked immunosorbent assay)-type procedure.

The lats gene can also be identified by mRNA

30 selection by nucleic acid hybridization followed by in vitro translation. In this procedure, fragments are used to isolate complementary mRNAs by hybridization. Such DNA fragments may represent available, purified lats DNA of another species (e.g., Drosophila, mouse, human).

35 Immunoprecipitation analysis or functional assays (e.g., aggregation ability in vitro; binding to receptor; see infra) of the in vitro translation products of the isolated products



of the isolated mRNAs identifies the mRNA and, therefore, the complementary DNA fragments that contain the desired sequences. In addition, specific mRNAs may be selected by adsorption of polysomes isolated from cells to immobilized antibodies specifically directed against lats protein. A radiolabelled lats cDNA can be synthesized using the selected mRNA (from the adsorbed polysomes) as a template. The radiolabelled mRNA or cDNA may then be used as a probe to identify the lats DNA fragments from among other genomic DNA fragments.

Alternatives to isolating the lats genomic DNA include, but are not limited to, chemically synthesizing the gene sequence itself from a known sequence or making cDNA to the mRNA which encodes the lats protein. For example, RNA for cDNA cloning of the lats gene can be isolated from cells which express lats. Other methods are possible and within the scope of the invention.

The identified and isolated gene can then be inserted into an appropriate cloning vector. A large number 20 of vector-host systems known in the art may be used. Possible vectors include, but are not limited to, plasmids or modified viruses, but the vector system must be compatible with the host cell used. Such vectors include, but are not limited to, bacteriophages such as lambda derivatives, or 25 plasmids such as PBR322 or pUC plasmid derivatives or the Bluescript vector (Stratagene). The insertion into a cloning vector can, for example, be accomplished by ligating the DNA fragment into a cloning vector which has complementary cohesive termini. However, if the complementary restriction 30 sites used to fragment the DNA are not present in the cloning vector, the ends of the DNA molecules may be enzymatically modified. Alternatively, any site desired may be produced by ligating nucleotide sequences (linkers) onto the DNA termini; these ligated linkers may comprise specific chemically 35 synthesized oligonucleotides encoding restriction endonuclease recognition sequences. In an alternative

method, the cleaved vector and lats gene may be modified by



homopolymeric tailing. Recombinant molecules can be introduced into host cells via transformation, transfection, infection, electroporation, etc., so that many copies of the gene sequence are generated.

- In an alternative method, the desired gene may be identified and isolated after insertion into a suitable cloning vector in a "shot gun" approach. Enrichment for the desired gene, for example, by size fractionization, can be done before insertion into the cloning vector.
- In specific embodiments, transformation of host cells with recombinant DNA molecules that incorporate the isolated lats gene, cDNA, or synthesized DNA sequence enables generation of multiple copies of the gene. Thus, the gene may be obtained in large quantities by growing transformants,
- 15 isolating the recombinant DNA molecules from the transformants and, when necessary, retrieving the inserted gene from the isolated recombinant DNA.

The lats sequences provided by the instant invention include those nucleotide sequences encoding

20 substantially the same amino acid sequences as found in native lats proteins, and those encoded amino acid sequences with functionally equivalent amino acids, as well as those encoding other lats derivatives or analogs, as described in Sections 5.6 and 5.6.1 infra for lats derivatives and

25 analogs.

#### 5.2. EXPRESSION OF THE LATS GENES

The nucleotide sequence coding for a lats protein or a functionally active analog or fragment or other

30 derivative thereof (see Section 5.6), can be inserted into an appropriate expression vector, i.e., a vector which contains the necessary elements for the transcription and translation of the inserted protein-coding sequence. The necessary transcriptional and translational signals can also be

35 supplied by the native lats gene and/or its flanking regions. A variety of host-vector systems may be utilized to express

the protein-coding sequence. These include but are not

limited to mammalian cell systems infected with virus (e.g., vaccinia virus, adenovirus, etc.); insect cell systems infected with virus (e.g., baculovirus); microorganisms such as yeast containing yeast vectors, or bacteria transformed with bacteriophage, DNA, plasmid DNA, or cosmid DNA. The expression elements of vectors vary in their strengths and specificities. Depending on the host-vector system utilized, any one of a number of suitable transcription and translation elements may be used. In specific embodiments, the human lats gene is expressed, or a sequence encoding a functionally active portion of human lats. In yet another embodiment, a fragment of lats comprising a domain of the lats protein is

Any of the methods previously described for the
insertion of DNA fragments into a vector may be used to
construct expression vectors containing a chimeric gene
consisting of appropriate transcriptional/translational
control signals and the protein coding sequences. These
methods may include in vitro recombinant DNA and synthetic
techniques and in vivo recombinants (genetic recombination).
Expression of nucleic acid sequence encoding a lats protein
or peptide fragment may be regulated by a second nucleic acid
sequence so that the lats protein or peptide is expressed in
a host transformed with the recombinant DNA molecule. For

- 25 example, expression of a lats protein may be controlled by any promoter/enhancer element known in the art. In a specific embodiment, the promoter is not a native lats gene promoter. Promoters which may be used to control lats expression include, but are not limited to, the SV40 early
- 30 promoter region (Bernoist and Chambon, 1981, Nature 290:304-310), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto, et al., 1980, Cell 22:787-797), the herpes thymidine kinase promoter (Wagner et al., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:1441-1445), the
- 35 regulatory sequences of the metallothionein gene (Brinster et al., 1982, Nature 296:39-42); prokaryotic expression vectors such as the  $\beta$ -lactamase promoter (Villa-Kamaroff, et al.,



1978, Proc. Natl. Acad. Sci. U.S.A. 75:3727-3731), or the tac promoter (DeBoer, et al., 1983, Proc. Natl. Acad. Sci. U.S.A. 80:21-25); see also "Useful proteins from recombinant bacteria" in Scientific American, 1980, 242:74-94; plant

- 5 expression vectors comprising the nopaline synthetase promoter region (Herrera-Estrella et al., Nature 303:209-213) or the cauliflower mosaic virus 35S RNA promoter (Gardner, et al., 1981, Nucl. Acids Res. 9:2871), and the promoter of the photosynthetic enzyme ribulose biphosphate carboxylase
- 10 (Herrera-Estrella et al., 1984, Nature 310:115-120); promoter elements from yeast or other fungi such as the Gal 4 promoter, the ADC (alcohol dehydrogenase) promoter, PGK (phosphoglycerol kinase) promoter, alkaline phosphatase promoter, and the following animal transcriptional control
- 15 regions, which exhibit tissue specificity and have been utilized in transgenic animals: elastase I gene control region which is active in pancreatic acinar cells (Swift et al., 1984, Cell 38:639-646; Ornitz et al., 1986, Cold Spring Harbor Symp. Quant. Biol. 50:399-409; MacDonald, 1987,
- 20 Hepatology 7:425-515); insulin gene control region which is active in pancreatic beta cells (Hanahan, 1985, Nature 315:115-122), immunoglobulin gene control region which is active in lymphoid cells (Grosschedl et al., 1984, Cell 38:647-658; Adames et al., 1985, Nature 318:533-538;
- 25 Alexander et al., 1987, Mol. Cell. Biol. 7:1436-1444), mouse mammary tumor virus control region which is active in testicular, breast, lymphoid and mast cells (Leder et al., 1986, Cell 45:485-495), albumin gene control region which is active in liver (Pinkert et al., 1987, Genes and Devel.
- 30 1:268-276), alpha-fetoprotein gene control region which is active in liver (Krumlauf et al., 1985, Mol. Cell. Biol. 5:1639-1648; Hammer et al., 1987, Science 235:53-58; alpha 1-antitrypsin gene control region which is active in the liver (Kelsey et al., 1987, Genes and Devel. 1:161-171), beta-
- 35 globin gene control region which is active in myeloid cells (Mogram et al., 1985, Nature 315:338-340; Kollias et al., 1986, Cell 46:89-94; myelin basic protein gene control region

which is active in oligodendrocyte cells in the brain (Readhead et al., 1987, Cell 48:703-712); myosin light chain2 gene control region which is active in skeletal muscle (Sani, 1985, Nature 314:283-286), and gonadotropic releasing hormone gene control region which is active in the hypothalamus (Mason et al., 1986, Science 234:1372-1378).

In a specific embodiment, a vector is used that comprises a promoter operably linked to a lats-encoding nucleic acid, one or more origins of replication, and,

10 optionally, one or more selectable markers (e.g., an antibiotic resistance gene).

In a specific embodiment, an expression construct is made by subcloning a lats coding sequence into the EcoRI restriction site of each of the three pGEX vectors

15 (Glutathione S-Transferase expression vectors; Smith and Johnson, 1988, Gene 7:31-40). This allows for the expression of the lats protein product from the subclone in the correct reading frame.

Expression vectors containing lats gene inserts can

20 be identified by three general approaches: (a) nucleic acid
hybridization, (b) presence or absence of "marker" gene
functions, and (c) expression of inserted sequences. In the
first approach, the presence of a lats gene inserted in an
expression vector can be detected by nucleic acid

- 25 hybridization using probes comprising sequences that are homologous to an inserted lats gene. In the second approach, the recombinant vector/host system can be identified and selected based upon the presence or absence of certain "marker" gene functions (e.g., thymidine kinase activity,
- 30 resistance to antibiotics, transformation phenotype, occlusion body formation in baculovirus, etc.) caused by the insertion of a lats gene in the vector. For example, if the lats gene is inserted within the marker gene sequence of the vector, recombinants containing the lats insert can be
- 35 identified by the absence of the marker gene function. In the third approach, recombinant expression vectors can be identified by assaying the lats product expressed by the



recombinant. Such assays can be based, for example, on the physical or functional properties of the lats protein in *in vitro* assay systems, e.g., kinase activity, binding with anti-lats antibody, inhibition of cell proliferation.

- Once a particular recombinant DNA molecule is identified and isolated, several methods known in the art may be used to propagate it. Once a suitable host system and growth conditions are established, recombinant expression vectors can be propagated and prepared in quantity. As
- 10 previously explained, the expression vectors which can be used include, but are not limited to, the following vectors or their derivatives: human or animal viruses such as vaccinia virus or adenovirus; insect viruses such as baculovirus; yeast vectors; bacteriophage vectors (e.g.,
- 15 lambda), and plasmid and cosmid DNA vectors, to name but a few.

In addition, a host cell strain may be chosen which modulates the expression of the inserted sequences, or modifies and processes the gene product in the specific

- 20 fashion desired. Expression from certain promoters can be elevated in the presence of certain inducers; thus, expression of the genetically engineered lats protein may be controlled. Furthermore, different host cells have characteristic and specific mechanisms for the translational
- 25 and post-translational processing and modification (e.g., glycosylation, phosphorylation of proteins. Appropriate cell lines or host systems can be chosen to ensure the desired modification and processing of the foreign protein expressed. For example, expression in a bacterial system can be used to
- 30 produce an unglycosylated core protein product. Expression in yeast will produce a glycosylated product. Expression in mammalian cells can be used to ensure "native" glycosylation of a heterologous protein. Furthermore, different vector/host expression systems may effect processing
- 35 reactions to different extents.

In other specific embodiments, the lats protein, fragment, analog, or derivative may be expressed as a fusion,

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or chimeric protein product (comprising the protein, fragment, analog, or derivative joined via a peptide bond to a heterologous protein sequence (of a different protein)). Such a chimeric product can be made by ligating the sappropriate nucleic acid sequences encoding the desired amino acid sequences to each other by methods known in the art, in the proper coding frame, and expressing the chimeric product by methods commonly known in the art. Alternatively, such a chimeric product may be made by protein synthetic techniques, e.g., by use of a peptide synthesizer.

Both cDNA and genomic sequences can be cloned and expressed.

# 5.3. IDENTIFICATION AND PURIFICATION OF THE LATS GENE PRODUCTS

In particular aspects, the invention provides amino acid sequences of lats, preferably human lats, and fragments and derivatives thereof which comprise an antigenic determinant (i.e., can be recognized by an antibody) or which are otherwise functionally active, as well as nucleic acid sequences encoding the foregoing. "Functionally active" lats material as used herein refers to that material displaying one or more known functional activities associated with a full-length (wild-type) lats protein, e.g., kinase activity, inhibition of cell proliferation, tumor inhibition, binding to an SH3-domain, binding to a lats substrate or lats binding partner, antigenicity (binding to an anti-lats antibody), immunogenicity, etc.

fragments of a lats protein consisting of at least 6 amino acids, 10 amino acids, 50 amino acids, or of at least 75 amino acids. In other embodiments, the proteins comprise or consist essentially of a lats carboxy (C)-terminal domain 3 (LCD3), lats C-terminal domain 2 (LCD2), lats C-terminal domain 1 (LCD1), kinase domain, kinase subdomains, lats flanking domain (amino-terminal to the kinase domain), lats split domain 1 (LSD1), lats split domain 2 (LSD2),



SH3-binding domain, and opa repeat domain (see Section 8 infra), or any combination of the foregoing, of a lats protein. Fragments, or proteins comprising fragments, lacking some or all of the foregoing regions of a lats protein are also provided. Nucleic acids encoding the foregoing are provided.

Once a recombinant which expresses the lats gene sequence is identified, the gene product can be analyzed. This is achieved by assays based on the physical or functional properties of the product, including radioactive labelling of the product followed by analysis by gel electrophoresis, immunoassay, etc.

Once the lats protein is identified, it may be isolated and purified by standard methods including

15 chromatography (e.g., ion exchange, affinity, and sizing column chromatography), centrifugation, differential solubility, or by any other standard technique for the purification of proteins. The functional properties may be evaluated using any suitable assay (see Section 5.7).

Alternatively, once a lats protein produced by a recombinant is identified, the amino acid sequence of the protein can be deduced from the nucleotide sequence of the chimeric gene contained in the recombinant. As a result, the protein can be synthesized by standard chemical methods known in the art (e.g., see Hunkapiller, M., et al., 1984, Nature 310:105-111).

In another alternate embodiment, native lats proteins can be purified from natural sources, by standard methods such as those described above (e.g., immunoaffinity 30 purification).

In a specific embodiment of the present invention, such lats proteins, whether produced by recombinant DNA techniques or by chemical synthetic methods or by purification of native proteins, include but are not limited 35 to those containing, as a primary amino acid sequence, all or part of the amino acid sequence substantially as depicted in Figure 9 (SEQ ID NO:4), as well as fragments and other

derivatives, and analogs thereof, including proteins homologous thereto.

5.4. STRUCTURE OF THE LATS GENE AND PROTEIN

The structure of the lats gene and protein can be analyzed by various methods known in the art.

#### 5.4.1. GENETIC ANALYSIS

The cloned DNA or cDNA corresponding to the lats

10 gene can be analyzed by methods including but not limited to Southern hybridization (Southern, E.M., 1975, J. Mol. Biol. 98:503-517), Northern hybridization (see e.g., Freeman et al., 1983, Proc. Natl. Acad. Sci. U.S.A. 80:4094-4098), restriction endonuclease mapping (Maniatis, T., 1982,

- York), and DNA sequence analysis. Polymerase chain reaction (PCR; U.S. Patent Nos. 4,683,202, 4,683,195 and 4,889,818; Gyllenstein et al., 1988, Proc. Natl. Acad. Sci. U.S.A. 85:7652-7656; Ochman et al., 1988, Genetics 120:621-623; Loh
- 20 et al., 1989, Science 243:217-220) followed by Southern hybridization with a lats-specific probe can allow the detection of the lats gene in DNA from various cell types. Methods of amplification other than PCR are commonly known and can also be employed. In one embodiment, Southern
- 25 hybridization can be used to determine the genetic linkage of lats. Northern hybridization analysis can be used to determine the expression of the lats gene. Various cell types, at various states of development or activity can be tested for lats expression. The stringency of the
- 30 hybridization conditions for both Southern and Northern hybridization can be manipulated to ensure detection of nucleic acids with the desired degree of relatedness to the specific lats probe used. Modifications of these methods and other methods commonly known in the art can be used.
- Restriction endonuclease mapping can be used to roughly determine the genetic structure of the lats gene.



Restriction maps derived by restriction endonuclease cleavage can be confirmed by DNA sequence analysis.

DNA sequence analysis can be performed by any techniques known in the art, including but not limited to the 5 method of Maxam and Gilbert (1980, Meth. Enzymol. 65:499-560), the Sanger dideoxy method (Sanger, F., et al., 1977, Proc. Natl. Acad. Sci. U.S.A. 74:5463), the use of T7 DNA polymerase (Tabor and Richardson, U.S. Patent No. 4,795,699), or use of an automated DNA sequenator (e.g., Applied Biosystems, Foster City, CA).

#### 5.4.2. PROTEIN ANALYSIS

The amino acid sequence of the lats protein can be derived by deduction from the DNA sequence, or alternatively, 15 by direct sequencing of the protein, e.g., with an automated amino acid sequencer.

The lats protein sequence can be further characterized by a hydrophilicity analysis (Hopp, T. and Woods, K., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:3824). A 20 hydrophilicity profile can be used to identify the hydrophobic and hydrophilic regions of the lats protein and the corresponding regions of the gene sequence which encode such regions.

Secondary, structural analysis (Chou, P. and 25 Fasman, G., 1974, Biochemistry 13:222) can also be done, to identify regions of lats that assume specific secondary structures.

Manipulation, translation, and secondary structure prediction, open reading frame prediction and plotting, as 30 well as determination of sequence homologies, can also be accomplished using computer software programs available in the art.

Other methods of structural analysis can also be employed. These include but are not limited to X-ray

35 crystallography (Engstom, A., 1974, Biochem. Exp. Biol. 11:713) and computer modeling (Fletterick, R. and Zoller, M.
(eds.), 1986, Computer Graphics and Molecular Modeling, in

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Current Communications in Molecular Biology, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York).

## 5.5. GENERATION OF ANTIBODIES TO LATS PROTEINS AND DERIVATIVES THEREOF

fragments or other derivatives, or analogs thereof, may be used as an immunogen to generate antibodies which immunospecifically bind such an immunogen. Such antibodies include but are not limited to polyclonal, monoclonal, chimeric, single chain, Fab fragments, and an Fab expression library. In a specific embodiment, antibodies to a human lats protein are produced. In another embodiment, antibodies to a domain (e.g., the SH3-binding domain) of a lats protein are produced. In a specific embodiment, fragments of a lats protein identified as hydrophilic are used as immunogens for antibody production.

Various procedures known in the art may be used for the production of polyclonal antibodies to a lats protein or 20 derivative or analog. In a particular embodiment, rabbit polyclonal antibodies to an epitope of a lats protein encoded by a sequence of SEQ ID NOS:2, 4, 6 or 8, or a subsequence thereof, can be obtained. For the production of antibody, various host animals can be immunized by injection with the native lats protein, or a synthetic version, or derivative (e.g., fragment) thereof, including but not limited to rabbits, mice, rats, etc. Various adjuvants may be used to increase the immunological response, depending on the host species, and including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanins, dinitrophenol, and potentially useful human adjuvants such as BCG (bacille Calmette-Guerin) and 35 corynebacterium parvum.

For preparation of monoclonal antibodies directed toward a lats protein sequence or analog thereof, any



technique which provides for the production of antibody molecules by continuous cell lines in culture may be used. For example, the hybridoma technique originally developed by Kohler and Milstein (1975, Nature 256:495-497), as well as

- 5 the trioma technique, the human B-cell hybridoma technique (Kozbor et al., 1983, Immunology Today 4:72), and the EBV-hybridoma technique to produce human monoclonal antibodies (Cole et al., 1985, in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). In an additional
- 10 embodiment of the invention, monoclonal antibodies can be produced in germ-free animals utilizing recent technology (PCT/US90/02545). According to the invention, human antibodies may be used and can be obtained by using human hybridomas (Cote et al., 1983, Proc. Natl. Acad. Sci. U.S.A.
- 15 80:2026-2030) or by transforming human B cells with EBV virus in vitro (Cole et al., 1985, in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, pp. 77-96). In fact, according to the invention, techniques developed for the production of "chimeric antibodies" (Morrison et al., 1984, Proc. Natl.
- 20 Acad. Sci. U.S.A. 81:6851-6855; Neuberger et al., 1984,
  Nature 312:604-608; Takeda et al., 1985, Nature 314:452-454)
  by splicing the genes from a mouse antibody molecule specific
  for lats together with genes from a human antibody molecule
  of appropriate biological activity can be used; such
- 25 antibodies are within the scope of this invention.

According to the invention, techniques described for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce lats-specific single chain antibodies. An additional embodiment of the invention

- 30 utilizes the techniques described for the construction of Fab expression libraries (Huse et al., 1989, Science 246:1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity for lats proteins, derivatives, or analogs.
- Antibody fragments which contain the idiotype of the molecule can be generated by known techniques. For example, such fragments include but are not limited to: the

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F(ab')<sub>2</sub> fragment which can be produced by pepsin digestion of the antibody molecule; the Fab' fragments which can be generated by reducing the disulfide bridges of the F(ab')<sub>2</sub> fragment, the Fab fragments which can be generated by treating the antibody molecule with papain and a reducing agent, and Fv fragments.

In the production of antibodies, screening for the desired antibody can be accomplished by techniques known in the art, e.g. ELISA (enzyme-linked immunosorbent assay). For example, to select antibodies which recognize a specific domain of a lats protein, one may assay generated hybridomas for a product which binds to a lats fragment containing such domain. For selection of an antibody that specifically binds a first lats homolog but which does not specifically bind a different lats homolog, one can select on the basis of positive binding to the first lats homolog and a lack of binding to the second lats homolog.

Antibodies specific to a domain of a lats protein are also provided.

20 The foregoing antibodies can be used in methods known in the art relating to the localization and activity of the lats protein sequences of the invention, e.g., for imaging these proteins, measuring levels thereof in appropriate physiological samples, in diagnostic methods, 25 etc.

In another embodiment of the invention (see infra), anti-lats antibodies and fragments thereof containing the binding domain are Therapeutics.

## 30 5.6. LATS PROTEINS, DERIVATIVES AND ANALOGS

The invention further relates to lats proteins, and derivatives (including but not limited to fragments) and analogs of lats proteins. Nucleic acids encoding lats protein derivatives and protein analogs are also provided.

35 In one embodiment, the lats proteins are encoded by the lats nucleic acids described in Section 5.1 supra. In particular aspects, the proteins, derivatives, or analogs are of lats



proteins of animals, e.g., fly, frog, mouse, rat, pig, cow, dog, monkey, human, or of plants.

The production and use of derivatives and analogs related to lats are within the scope of the present

- 5 invention. In a specific embodiment, the derivative or analog is functionally active, i.e., capable of exhibiting one or more functional activities associated with a full-length, wild-type lats protein. As one example, such derivatives or analogs which have the desired immunogenicity
- 10 or antigenicity can be used, for example, in immunoassays, for immunization, for inhibition of lats activity, etc. As another example, such derivatives or analogs which have the desired kinase activity, or which are phosphorylated or dephosphorylated, are provided. Derivatives or analogs that
- 15 retain, or alternatively lack or inhibit, a desired lats property of interest (e.g., binding to an SH3-domain-containing protein or other lats binding partner, kinase activity, inhibition of cell proliferation, tumor inhibition), can be used as inducers, or inhibitors,
- 20 respectively, of such property and its physiological correlates. A specific embodiment relates to a lats fragment that can be bound by an anti-lats antibody. Derivatives or analogs of lats can be tested for the desired activity by procedures known in the art, including but not limited to the 25 assays described in Sections 5.7 and 5.9.

In particular, lats derivatives can be made by altering lats sequences by substitutions, additions or deletions that provide for functionally equivalent molecules. Due to the degeneracy of nucleotide coding sequences, other

- 30 DNA sequences which encode substantially the same amino acid sequence as a lats gene may be used in the practice of the present invention. These include but are not limited to nucleotide sequences comprising all or portions of lats genes which are altered by the substitution of different codons
- 35 that encode a functionally equivalent amino acid residue within the sequence, thus producing a silent change. Likewise, the lats derivatives of the invention include, but

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are not limited to, those containing, as a primary amino acid sequence, all or part of the amino acid sequence of a lats protein including altered sequences in which functionally equivalent amino acid residues are substituted for residues 5 within the sequence resulting in a silent change. For example, one or more amino acid residues within the sequence can be substituted by another amino acid of a similar polarity which acts as a functional equivalent, resulting in a silent alteration. Substitutes for an amino acid within 10 the sequence may be selected from other members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, 15 serine, threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid.

- In a specific embodiment of the invention, proteins consisting of or comprising a fragment of a lats protein consisting of at least 10 (continuous) amino acids of the lats protein is provided. In other embodiments, the fragment consists of at least 20 or 50 amino acids of the lats
- 25 protein. In specific embodiments, such fragments are not larger than 35, 100 or 200 amino acids. Derivatives or analogs of lats include but are not limited to those molecules comprising regions that are substantially homologous to lats or fragments thereof (e.g., in various
- 30 embodiments, at least 60% or 70% or 80% or 90% or 95% identity over an amino acid sequence of identical size or when compared to an aligned sequence in which the alignment is done by a computer homology program known in the art) or whose encoding nucleic acid is capable of hybridizing to a
- 35 coding lats sequence, under stringent, moderately stringent, or nonstringent conditions.

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The lats derivatives and analogs of the invention can be produced by various methods known in the art. The manipulations which result in their production can occur at

- the gene or protein level. For example, the cloned lats gene
  5 sequence can be modified by any of numerous strategies known in the art (Maniatis, T., 1990, Molecular Cloning, A Laboratory Manual, 2d ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, New York). The sequence can be cleaved at appropriate sites with restriction endonuclease(s),
- 10 followed by further enzymatic modification if desired, isolated, and ligated in vitro. In the production of the gene encoding a derivative or analog of lats, care should be taken to ensure that the modified gene remains within the same translational reading frame as lats, uninterrupted by
- 15 translational stop signals, in the gene region where the desired lats activity is encoded.

Additionally, the lats-encoding nucleic acid sequence can be mutated in vitro or in vivo, to create and/or destroy translation, initiation, and/or termination

- 20 sequences, or to create variations in coding regions and/or form new restriction endonuclease sites or destroy preexisting ones, to facilitate further in vitro modification. Any technique for mutagenesis known in the art can be used, including but not limited to, chemical
- 25 mutagenesis, in vitro site-directed mutagenesis (Hutchinson,
  C., et al., 1978, J. Biol. Chem 253:6551), use of TAB®
  linkers (Pharmacia), etc.

Manipulations of the lats sequence may also be made at the protein level. Included within the scope of the 30 invention are lats protein fragments or other derivatives or

- analogs which are differentially modified during or after translation, e.g., by glycosylation, acetylation, phosphorylation, amidation, derivatization by known protecting/blocking groups, proteolytic cleavage, linkage to
- 35 an antibody molecule or other cellular ligand, etc. Any of numerous chemical modifications may be carried out by known techniques, including but not limited to specific chemical



cleavage by cyanogen bromide, trypsin, chymotrypsin, papain, V8 protease, NaBH4; acetylation, formylation, oxidation, reduction; metabolic synthesis in the presence of tunicamycin; etc.

- In addition, analogs and derivatives of lats can be chemically synthesized. For example, a peptide corresponding to a portion of a lats protein which comprises the desired domain (see Section 5.6.1), or which mediates the desired activity in vitro, can be synthesized by use of a peptide
- 10 synthesizer. Furthermore, if desired, nonclassical amino acids or chemical amino acid analogs can be introduced as a substitution or addition into the lats sequence. Nonclassical amino acids include but are not limited to the Disomers of the common amino acids, α-amino isobutyric acid,
- 15 4-aminobutyric acid, Abu, 2-amino butyric acid, γ-Abu, ε-Ahx, 6-amino hexanoic acid, Aib, 2-amino isobutyric acid, 3-amino propionic acid, ornithine, norleucine, norvaline, hydroxyproline, sarcosine, citrulline, cysteic acid, t-butylglycine, t-butylalanine, phenylglycine,
- 20 cyclohexylalanine,  $\beta$ -alanine, fluoro-amino acids, designer amino acids such as  $\beta$ -methyl amino acids,  $C\alpha$ -methyl amino acids,  $N\alpha$ -methyl amino acids, and amino acid analogs in general. Furthermore, the amino acid can be D (dextrorotary) or L (levorotary).
- In a specific embodiment, the lats derivative is a chimeric, or fusion, protein comprising a lats protein or fragment thereof (preferably consisting of at least a domain or motif of the lats protein, or at least 10 amino acids of the lats protein) joined at its amino- or carboxy-terminus
- 30 via a peptide bond to an amino acid sequence of a different protein. In one embodiment, such a chimeric protein is produced by recombinant expression of a nucleic acid encoding the protein (comprising a lats-coding sequence joined inframe to a coding sequence for a different protein). Such a
- 35 chimeric product can be made by ligating the appropriate nucleic acid sequences encoding the desired amino acid sequences to each other by methods known in the art, in the

proper coding frame, and expressing the chimeric product by methods commonly known in the art. Alternatively, such a chimeric product may be made by protein synthetic techniques, e.g., by use of a peptide synthesizer. Chimeric genes comprising portions of lats fused to any heterologous protein-encoding sequences may be constructed. A specific embodiment relates to a chimeric protein comprising a fragment of lats of at least six amino acids.

In another specific embodiment, the lats derivative

10 is a molecule comprising a region of homology with a lats
protein. By way of example, in various embodiments, a first
protein region can be considered "homologous" to a second
protein region when the amino acid sequence of the first
region is at least 30%, 40%, 50%, 60%, 70%, 75%, 80%, 90%, or

15 95% identical, when compared to any sequence in the second
region of an equal number of amino acids as the number
contained in the first region or when compared to an aligned
sequence of the second region that has been aligned by a
computer homology program known in the art. For example, a

20 molecule can comprise one or more regions homologous to a
lats domain (see Section 5.6.1) or a portion thereof.

Other specific embodiments of derivatives and analogs are described in the subsection below and examples sections infra.

25

## 5.6.1. DERIVATIVES OF LATS CONTAINING ONE OR MORE DOMAINS OF THE PROTEIN

In a specific embodiment, the invention relates to lats derivatives and analogs, in particular lats fragments and derivatives of such fragments, that comprise, or alternatively consist of, one or more domains of a lats protein, including but not limited to a lats C-terminal domain 3 (LCD3), lats C-terminal domain 2 (LCD2), lats C-terminal domain 1 (LCD1), kinase domain, kinase subdomains, lats flanking domain (LFD) (amino-terminal to the kinase domain), lats split domain 1 (LSD1), lats split domain 2 (LSD2), SH3-binding domain, and opa repeat domain, functional

(e.g., binding) fragments of any of the foregoing, or any combination of the foregoing. In particular examples relating to the human, mouse and Drosophila lats proteins, such domains are identified in Examples Sections 6 and 8, and 5 in Figures 6A, 6B, and 13.

A specific embodiment relates to molecules comprising specific fragments of lats that are those fragments in the respective lats protein most homologous to specific fragments of a human or mouse lats protein. A 10 fragment comprising a domain of a lats homolog can be identified by protein analysis methods as described in Sections 5.3.2 or 6.

In a specific embodiment, a lats protein, derivative or analog is provided that has a kinase domain and 15 has a phosphorylated serine situated within 20 residues upstream of an Ala-Pro-Glu consensus in subdomain eight of its kinase domain. In another embodiment, a lats protein derivative or analog is provided with a kinase domain and with a dephosphorylated serine situated within 20 residues 20 upstream of an Ala-Pro-Glu consensus in subdomain eight of its kinase domain, or in which the serine situated within 20 residues upstream of that consensus has been deleted or substituted by another amino acid. In a specific embodiment, the invention provides various phosphorylated and 25 dephosphorylated forms of the lats protein, derivative, or analog that are active kinase forms. Both phosphorylation and dephosphorylation of lats at different residues could potentially activate lats. In another specific embodiment, the invention provides various phosphorylated and 30 dephosphorylated forms of the lats protein, derivative or analog that are inactive kinase forms. Phosphorylation can be carried out by any methods known in the art, e.g., by use of a kinase. Dephosphorylation can be carried out by use of

another specific embodiment relates to a derivative or analog of a lats protein that is a dominant-active protein kinase. Such a derivative or analog comprises a lats kinase

any methods known in the art, e.g., by use of a phosphatase.

dominant active lats kinase.

- Another specific embodiment relates to a derivative or analog of lats that is a dominant-negative protein kinase. Protein kinases can be mutated into dominant negative forms. Expression of a dominant negative protein kinase can suppress the activity of the wild-type form of the same kinase.
- 20 Dominant negative forms of protein kinases are often obtained by expressing an inactive form of a kinase (Milarski and Saltiel, 1994, J. Biol. Chem. 269(33):21239-21243) or by expressing a noncatalytic domain of a kinase (Lu and Means, 1994, EMBO J. 12:2103-2113; Yarden et al., 1992, EMBO J.
- 25 11:2159-2166). Thus, a lats dominant-negative kinase can be obtained by mutating the kinase domain so as to be inactive (e.g., by deletion and/or point mutation). By way of example, a lats derivative that is a dominant-negative kinase is a lats protein that lacks a kinase domain but comprises
- 30 one or more of the other domains of the lats protein; e.g., a lats protein derivative truncated at about the beginning of the kinase domain (i.e., a lats fragment containing only sequences amino-terminal to the kinase domain). By way of another example, a lats derivative that is a dominant-
- 35 negative kinase is a lats protein in which one of the residues conserved among serine/threonine kinases (see Hanks

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et al., 1988, Science 241:42-52) is mutated (deleted or substituted by a different residue).

In another specific embodiment, a molecule is provided that comprises one or more domains (or functional 5 portion thereof) of a lats protein but that also lacks one or more domains (or functional portion thereof) of a lats protein. In particular examples, lats protein derivatives are provided that lack an opa repeat domain. By way of another example, such a protein may also lack all or a 10 portion of the kinase domain, but retain at least the SH3-binding domain of a lats protein. In another embodiment, a molecule is provided that comprises one or more domains (or functional portion thereof) of a lats protein, and that has one or more mutant (e.g., due to deletion or point 15 mutation(s)) domains of a lats protein (e.g., such that the mutant domain has decreased function). By way of example, the kinase domain may be mutant so as to have reduced, absent, or increased kinase activity.

### 5.7. ASSAYS OF LATS PROTEINS, <u>DERIVATIVES AND ANALOGS</u>

The functional activity of lats proteins, derivatives and analogs can be assayed by various methods.

assaying for the ability to bind or compete with wild-type lats for binding to anti-lats antibody, various immunoassays known in the art can be used, including but not limited to competitive and non-competitive assay systems using techniques such as radioimmunoassays, ELISA (enzyme linked immunosorbent assay), "sandwich" immunoassays, immunoradiometric assays, gel diffusion precipitin reactions, immunodiffusion assays, in situ immunoassays (using colloidal gold, enzyme or radioisotope labels, for example), western blots, precipitation reactions, agglutination assays (e.g.,

gel agglutination assays, hemagglutination assays), complement fixation assays, immunofluorescence assays, protein A assays, and immunoelectrophoresis assays, etc. In one embodiment, antibody binding is detected by detecting a label on the primary antibody. In another embodiment, the primary antibody is detected by detecting binding of a secondary antibody or reagent to the primary antibody. In a further embodiment, the secondary antibody is labelled. Many means are known in the art for detecting binding in an immunoassay and are within the scope of the present invention.

In another embodiment, where a lats-binding protein 10 is identified, the binding can be assayed, e.g., by means well-known in the art. In another embodiment, physiological correlates of lats binding to its substrates (signal transduction) can be assayed.

In another embodiment, kinase assays can be used to

15 measure lats kinase activity. Such assays can be carried out
by methods well known in the art. By way of example, a lats
protein is contacted with a substrate (e.g., a known
substrate of serine/threonine kinases) in the presence of a

17P-labeled phosphate donor, and any phosphorylation of the

20 substrate is detected or measured.

In another embodiment, in insect or other model systems, genetic studies can be done to study the phenotypic effect of a lats mutant that is a derivative or analog of wild-type lats (see Section 6, infra).

In addition, assays that can be used to detect or measure the ability to inhibit, or alternatively promote, cell proliferation are described in Section 5.9.

Other methods will be known to the skilled artisan and are within the scope of the invention.

30

### 5.8. THERAPEUTIC USES

The invention provides for treatment or prevention of various diseases and disorders by administration of a therapeutic compound (termed herein "Therapeutic"). Such "Therapeutics" include but are not limited to: lats proteins and analogs and derivatives (including fragments) thereof (e.g., as described hereinabove); antibodies thereto (as

described hereinabove); nucleic acids encoding the lats proteins, analogs, or derivatives (e.g., as described hereinabove); lats antisense nucleic acids, and lats agonists and antagonists. Disorders involving cell overproliferation are treated or prevented by administration of a Therapeutic that promotes lats function. Disorders in which cell proliferation is deficient or is desired are treated or prevented by administration of a Therapeutic that antagonizes (inhibits) lats function. The above is described in detail in the subsections below.

Generally, administration of products of a species origin or species reactivity (in the case of antibodies) that is the same species as that of the patient is preferred. Thus, in a preferred embodiment, a human lats protein, 15 derivative, or analog, or nucleic acid, or an antibody to a human lats protein, is therapeutically or prophylactically administered to a human patient.

Additional descriptions and sources of Therapeutics that can be used according to the invention are found in 20 Sections 5.1 through 5.7 herein.

# 5.8.1. TREATMENT AND PREVENTION OF DISORDERS INVOLVING OVERPROLIFERATION OF CELLS

overproliferation are treated or prevented by administration of a Therapeutic that promotes (i.e., increases or supplies) lats function. Examples of such a Therapeutic include but are not limited to lats proteins, derivatives, or fragments that are functionally active, particularly that are active in inhibiting cell proliferation (e.g., as demonstrated in in vitro assays or in animal models or in Drosophila), and nucleic acids encoding a lats protein or functionally active derivative or fragment thereof (e.g., for use in gene therapy). Other Therapeutics that can be used, e.g., lats agonists, can be identified using in vitro assays or animal models, or assays in Drosophila, examples of which are described infra.

In specific embodiments, Therapeutics that promote lats function are administered therapeutically (including prophylactically): (1) in diseases or disorders involving an absence or decreased (relative to normal or desired) level of

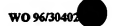
- 5 lats protein or function, for example, in patients where lats protein is lacking, genetically defective, biologically inactive or underactive, or underexpressed; or (2) in diseases or disorders wherein in vitro (or in vivo) assays (see infra) indicate the utility of lats agonist
- 10 administration. The absence or decreased level in lats protein or function can be readily detected, e.g., by obtaining a patient tissue sample (e.g., from biopsy tissue) and assaying it in vitro for RNA or protein levels, structure and/or activity of the expressed lats RNA or protein. Many
- 15 methods standard in the art can be thus employed, including but not limited to kinase assays, immunoassays to detect and/or visualize lats protein (e.g., Western blot, immunoprecipitation followed by sodium dodecyl sulfate polyacrylamide gel electrophoresis, immunocytochemistry,
- 20 etc.) and/or hybridization assays to detect lats expression by detecting and/or visualizing lats mRNA (e.g., Northern assays, dot blots, in situ hybridization, etc.), etc.

Diseases and disorders involving cell
overproliferation that can be treated or prevented include
25 but are not limited to malignancies, premalignant conditions
(e.g., hyperplasia, metaplasia, dysplasia), benign tumors,
hyperproliferative disorders, benign dysproliferative
disorders, etc. Examples of these are detailed below.

In a specific embodiment, the Therapeutic used,

30 that promotes lats function, is a lats protein, derivative or
analog comprising a lats kinase domain (and optionally also a
lats LFD, or the remainder of the lats sequence) in which a
serine within 20 residues upstream of the Ala-Pro-Glu
consensus in subdomain eight of the kinase domain is

35 phosphorylated or substituted by another residue (e.g., Glu,
Asp).



In another specific embodiment, the Therapeutic used, that promotes lats function, is a derivative or analog comprising a kinase domain of a lats protein that has been mutated so as to be dominantly active.

5.8.1.1. MALIGNANCIES

Malignancies and related disorders that can be treated or prevented by administration of a Therapeutic that promotes lats function include but are not limited to those 10 listed in Table 1 (for a review of such disorders, see Fishman et al., 1985, Medicine, 2d Ed., J.B. Lippincott Co., Philadelphia):

#### TABLE 1 MALIGNANCIES AND RELATED DISORDERS

Leukemia

acute leukemia

acute lymphocytic leukemia acute myelocytic leukemia

20

15

5

myeloblastic promyelocytic myelomonocytic monocytic

erythroleukemia

chronic leukemia

chronic myelocytic (granulocytic) leukemia chronic lymphocytic leukemia

25 Polycythemia vera

Lymphoma

Hodgkin's disease non-Hodgkin's disease

Multiple myeloma

Waldenström's macroglobulinemia

Heavy chain disease

Solid tumors 30

sarcomas and carcinomas

fibrosarcoma myxosarcoma liposarcoma chondrosarcoma osteogenic sarcoma

chordoma 35 angiosarcoma

endotheliosarcoma lymphangiosarcoma

35

lymphangioendotheliosarcoma synovioma mesothelioma Ewing's tumor leiomyosarcoma rhabdomyosarcoma 5 colon carcinoma pancreatic cancer breast cancer ovarian cancer prostate cancer squamous cell carcinoma basal cell carcinoma adenocarcinoma 10 sweat gland carcinoma sebaceous gland carcinoma papillary carcinoma papillary adenocarcinomas cystadenocarcinoma medullary carcinoma bronchogenic carcinoma 15 renal cell carcinoma hepatoma bile duct carcinoma choriocarcinoma seminoma embryonal carcinoma Wilms' tumor cervical cancer 20 uterine cancer testicular tumor lung carcinoma small cell lung carcinoma bladder carcinoma epithelial carcinoma glioma 25 astrocytoma medulloblastoma craniopharyngioma ependymoma pinealoma hemangioblastoma acoustic neuroma oligodendroglioma 30 menangioma melanoma neuroblastoma retinoblastoma

In specific embodiments, malignancy or dysproliferative changes (such as metaplasias and



gall bladder.

dysplasias), or hyperproliferative disorders, are treated or prevented in the bladder, breast, colon, lung, melanoma, pancreas, or uterus. In other specific embodiments, sarcoma, or leukemia is treated or prevented.

5.8.1.2. PREMALIGNANT CONDITIONS

The Therapeutics of the invention that promote lats activity can also be administered to treat premalignant conditions and to prevent progression to a neoplastic or 10 malignant state, including but not limited to those disorders listed in Table 1. Such prophylactic or therapeutic use is indicated in conditions known or suspected of preceding progression to neoplasia or cancer, in particular, where nonneoplastic cell growth consisting of hyperplasia, metaplasia, 15 or most particularly, dysplasia has occurred (for review of such abnormal growth conditions, see Robbins and Angell, 1976, Basic Pathology, 2d Ed., W.B. Saunders Co., Philadelphia, pp. 68-79.) Hyperplasia is a form of controlled cell proliferation involving an increase in cell 20 number in a tissue or organ, without significant alteration in structure or function. As but one example, endometrial hyperplasia often precedes endometrial cancer. Metaplasia is a form of controlled cell growth in which one type of adult or fully differentiated cell substitutes for another type of 25 adult cell. Metaplasia can occur in epithelial or connective tissue cells. Atypical metaplasia involves a somewhat disorderly metaplastic epithelium. Dysplasia is frequently a forerunner of cancer, and is found mainly in the epithelia; it is the most disorderly form of non-neoplastic cell growth, 30 involving a loss in individual cell uniformity and in the architectural orientation of cells. Dysplastic cells often have abnormally large, deeply stained nuclei, and exhibit pleomorphism. Dysplasia characteristically occurs where there exists chronic irritation or inflammation, and is often 35 found in the cervix, respiratory passages, oral cavity, and

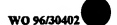


Alternatively or in addition to the presence of abnormal cell growth characterized as hyperplasia, metaplasia, or dysplasia, the presence of one or more characteristics of a transformed phenotype, or of a malignant 5 phenotype, displayed in vivo or displayed in vitro by a cell sample from a patient, can indicate the desirability of prophylactic/therapeutic administration of a Therapeutic that promotes lats function. As mentioned supra, such characteristics of a transformed phenotype include morphology 10 changes, looser substratum attachment, loss of contact inhibition, loss of anchorage dependence, protease release, increased sugar transport, decreased serum requirement, expression of fetal antigens, disappearance of the 250,000 dalton cell surface protein, etc. (see also id., at pp. 84-90 15 for characteristics associated with a transformed or malignant phenotype).

In a specific embodiment, leukoplakia, a benignappearing hyperplastic or dysplastic lesion of the
epithelium, or Bowen's disease, a carcinoma in situ, are pre20 neoplastic lesions indicative of the desirability of
prophylactic intervention.

In another embodiment, fibrocystic disease (cystic hyperplasia, mammary dysplasia, particularly adenosis (benign epithelial hyperplasia)) is indicative of the desirability of prophylactic intervention.

In other embodiments, a patient which exhibits one or more of the following predisposing factors for malignancy is treated by administration of an effective amount of a Therapeutic: a chromosomal translocation associated with a malignancy (e.g., the Philadelphia chromosome for chronic myelogenous leukemia, t(14;18) for follicular lymphoma, etc.), familial polyposis or Gardner's syndrome (possible forerunners of colon cancer), benign monoclonal gammopathy (a possible forerunner of multiple myeloma), and a first degree kinship with persons having a cancer or precancerous disease showing a Mendelian (genetic) inheritance pattern (e.g., familial polyposis of the colon, Gardner's syndrome,



hereditary exostosis, polyendocrine adenomatosis, medullary thyroid carcinoma with amyloid production and pheochromocytoma, Peutz-Jeghers syndrome, neurofibromatosis of Von Recklinghausen, retinoblastoma, carotid body tumor, cutaneous melanocarcinoma, intraocular melanocarcinoma, xeroderma pigmentosum, ataxia telangiectasia, Chediak-Higashi syndrome, albinism, Fanconi's aplastic anemia, and Bloom's syndrome; see Robbins and Angell, 1976, Basic Pathology, 2d Ed., W.B. Saunders Co., Philadelphia, pp. 112-113) etc.)

In another specific embodiment, a Therapeutic of the invention is administered to a human patient to prevent progression to breast, colon, lung, pancreatic, or uterine cancer, or melanoma or sarcoma.

#### 5.8.1.3. HYPERPROLIFERATIVE AND DYSPROLIFERATIVE DISORDERS

In another embodiment of the invention, a
Therapeutic that promotes lats activity is used to treat or
prevent hyperproliferative or benign dysproliferative

disorders. Specific embodiments are directed to treatment or
prevention of cirrhosis of the liver (a condition in which
scarring has overtaken normal liver regeneration processes),
treatment of keloid (hypertrophic scar) formation
(disfiguring of the skin in which the scarring process
interferes with normal renewal), psoriasis (a common skin
condition characterized by excessive proliferation of the
skin and delay in proper cell fate determination), benign
tumors, fibrocystic conditions, and tissue hypertrophy (e.g.,
prostatic hyperplasia).

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### 5.8.1.4. GENE THERAPY

In a specific embodiment, nucleic acids comprising a sequence encoding a lats protein or functional derivative thereof, are administered to promote lats function, by way of gene therapy. Gene therapy refers to therapy performed by the administration of a nucleic acid to a subject. In this embodiment of the invention, the nucleic acid produces its



encoded protein that mediates a therapeutic effect by promoting lats function.

Any of the methods for gene therapy available in the art can be used according to the present invention.

5 Exemplary methods are described below.

For general reviews of the methods of gene therapy, see Goldspiel et al., 1993, Clinical Pharmacy 12:488-505; Wu and Wu, 1991, Biotherapy 3:87-95; Tolstoshev, 1993, Ann. Rev. Pharmacol. Toxicol. 32:573-596; Mulligan, 1993, Science

- 10 260:926-932; and Morgan and Anderson, 1993, Ann. Rev. Biochem. 62:191-217; May, 1993, TIBTECH 11(5):155-215). Methods commonly known in the art of recombinant DNA technology which can be used are described in Ausubel et al. (eds.), 1993, Current Protocols in Molecular Biology, John
- 15 Wiley & Sons, NY; and Kriegler, 1990, Gene Transfer and Expression, A Laboratory Manual, Stockton Press, NY.

In a preferred aspect, the Therapeutic comprises a lats nucleic acid that is part of an expression vector that expresses a lats protein or fragment or chimeric protein

- 20 thereof in a suitable host. In particular, such a nucleic acid has a promoter operably linked to the lats coding region, said promoter being inducible or constitutive, and, optionally, tissue-specific. In another particular embodiment, a nucleic acid molecule is used in which the lats
- 25 coding sequences and any other desired sequences are flanked by regions that promote homologous recombination at a desired site in the genome, thus providing for intrachromosomal expression of the lats nucleic acid (Koller and Smithies, 1989, Proc. Natl. Acad. Sci. USA 86:8932-8935; Zijlstra et
- 30 al., 1989, Nature 342:435-438).

Delivery of the nucleic acid into a patient may be either direct, in which case the patient is directly exposed to the nucleic acid or nucleic acid-carrying vector, or indirect, in which case, cells are first transformed with the nucleic acid in vitro, then transplanted into the patient. These two approaches are known, respectively, as in vivo or ex vivo gene therapy.

In a specific embodiment, the nucleic acid is directly administered in vivo, where it is expressed to produce the encoded product. This can be accomplished by any of numerous methods known in the art, e.g., by constructing

- 5 it as part of an appropriate nucleic acid expression vector and administering it so that it becomes intracellular, e.g., by infection using a defective or attenuated retroviral or other viral vector (see U.S. Patent No. 4,980,286), or by direct injection of naked DNA, or by use of microparticle
- 10 bombardment (e.g., a gene gun; Biolistic, Dupont), or coating with lipids or cell-surface receptors or transfecting agents, encapsulation in liposomes, microparticles, or microcapsules, or by administering it in linkage to a peptide which is known to enter the nucleus, by administering it in linkage to a
- 15 ligand subject to receptor-mediated endocytosis (see e.g., Wu and Wu, 1987, J. Biol. Chem. 262:4429-4432) (which can be used to target cell types specifically expressing the receptors), etc. In another embodiment, a nucleic acidligand complex can be formed in which the ligand comprises a
- 20 fusogenic viral peptide to disrupt endosomes, allowing the nucleic acid to avoid lysosomal degradation. In yet another embodiment, the nucleic acid can be targeted in vivo for cell specific uptake and expression, by targeting a specific receptor (see, e.g., PCT Publications WO 92/06180 dated April
- 25 16, 1992 (Wu et al.); WO 92/22635 dated December 23, 1992 (Wilson et al.); WO92/20316 dated November 26, 1992 (Findeis et al.); WO93/14188 dated July 22, 1993 (Clarke et al.), WO 93/20221 dated October 14, 1993 (Young)). Alternatively, the nucleic acid can be introduced intracellularly and
- 30 incorporated within host cell DNA for expression, by homologous recombination (Koller and Smithies, 1989, Proc. Natl. Acad. Sci. USA 86:8932-8935; Zijlstra et al., 1989, Nature 342:435-438).

In a specific embodiment, a viral vector that

35 contains the lats nucleic acid is used. For example, a
retroviral vector can be used (see Miller et al., 1993, Meth.
Enzymol. 217:581-599). These retroviral vectors have been



modified to delete retroviral sequences that are not necessary for packaging of the viral genome and integration into host cell DNA. The lats nucleic acid to be used in gene therapy is cloned into the vector, which facilitates delivery

- 5 of the gene into a patient. More detail about retroviral vectors can be found in Boesen et al., 1994, Biotherapy 6:291-302, which describes the use of a retroviral vector to deliver the mdrl gene to hematopoietic stem cells in order to make the stem cells more resistant to chemotherapy. Other
- 10 references illustrating the use of retroviral vectors in gene therapy are: Clowes et al., 1994, J. Clin. Invest. 93:644-651; Kiem et al., 1994, Blood 83:1467-1473; Salmons and Gunzberg, 1993, Human Gene Therapy 4:129-141; and Grossman and Wilson, 1993, Curr. Opin. in Genetics and Devel. 3:110-15 114.

Adenoviruses are other viral vectors that can be used in gene therapy. Adenoviruses are especially attractive vehicles for delivering genes to respiratory epithelia. Adenoviruses naturally infect respiratory epithelia where

- 20 they cause a mild disease. Other targets for adenovirus-based delivery systems are liver, the central nervous system, endothelial cells, and muscle. Adenoviruses have the advantage of being capable of infecting non-dividing cells. Kozarsky and Wilson, 1993, Current Opinion in Genetics and
- 25 Development 3:499-503 present a review of adenovirus-based gene therapy. Bout et al., 1994, Human Gene Therapy 5:3-10 demonstrated the use of adenovirus vectors to transfer genes to the respiratory epithelia of rhesus monkeys. Other instances of the use of adenoviruses in gene therapy can be
- 30 found in Rosenfeld et al., 1991, Science 252:431-434;
  Rosenfeld et al., 1992, Cell 68:143-155; and Mastrangeli et
  al., 1993, J. Clin. Invest. 91:225-234.

Adeno-associated virus (AAV) has also been proposed for use in gene therapy (Walsh et al., 1993, Proc. Soc. Exp. 35 Biol. Med. 204:289-300.

Another approach to gene therapy involves transferring a gene to cells in tissue culture by such

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methods as electroporation, lipofection, calcium phosphate mediated transfection, or viral infection. Usually, the method of transfer includes the transfer of a selectable marker to the cells. The cells are then placed under selection to isolate those cells that have taken up and are expressing the transferred gene. Those cells are then delivered to a patient.

In this embodiment, the nucleic acid is introduced into a cell prior to administration in vivo of the resulting 10 recombinant cell. Such introduction can be carried out by any method known in the art, including but not limited to transfection, electroporation, microinjection, infection with a viral or bacteriophage vector containing the nucleic acid sequences, cell fusion, chromosome-mediated gene transfer,

- Numerous techniques are known in the art for the introduction of foreign genes into cells (see e.g., Loeffler and Behr, 1993, Meth. Enzymol. 217:599-618; Cohen et al., 1993, Meth. Enzymol. 217:618-644; Cline, 1985, Pharmac. Ther. 29:69-92)
- 20 and may be used in accordance with the present invention, provided that the necessary developmental and physiological functions of the recipient cells are not disrupted. The technique should provide for the stable transfer of the nucleic acid to the cell, so that the nucleic acid is
- 25 expressible by the cell and preferably heritable and expressible by its cell progeny.

The resulting recombinant cells can be delivered to a patient by various methods known in the art. In a preferred embodiment, epithelial cells are injected, e.g., subcutaneously. In another embodiment, recombinant skin cells may be applied as a skin graft onto the patient. Recombinant blood cells (e.g., hematopoietic stem or progenitor cells) are preferably administered intravenously. The amount of cells envisioned for use depends on the desired

35 effect, patient state, etc., and can be determined by one skilled in the art.



Cells into which a nucleic acid can be introduced for purposes of gene therapy encompass any desired, available cell type, and include but are not limited to epithelial cells, endothelial cells, keratinocytes, fibroblasts, muscle cells, hepatocytes; blood cells such as T lymphocytes, B lymphocytes, monocytes, macrophages, neutrophils, eosinophils, megakaryocytes, granulocytes; various stem or progenitor cells, in particular hematopoietic stem or progenitor cells, e.g., as obtained from bone marrow, umbilical cord blood, peripheral blood, fetal liver, etc.

In a preferred embodiment, the cell used for gene therapy is autologous to the patient.

In an embodiment in which recombinant cells are used in gene therapy, a lats nucleic acid is introduced into 15 the cells such that it is expressible by the cells or their progeny, and the recombinant cells are then administered in vivo for therapeutic effect. In a specific embodiment, stem or progenitor cells are used. Any stem and/or progenitor cells which can be isolated and maintained in vitro can

- 20 potentially be used in accordance with this embodiment of the present invention. Such stem cells include but are not limited to hematopoietic stem cells (HSC), stem cells of epithelial tissues such as the skin and the lining of the gut, embryonic heart muscle cells, liver stem cells (PCT
- 25 Publication WO 94/08598, dated April 28, 1994), and neural stem cells (Stemple and Anderson, 1992, Cell 71:973-985).

Epithelial stem cells (ESCs) or keratinocytes can be obtained from tissues such as the skin and the lining of the gut by known procedures (Rheinwald, 1980, Meth. Cell Bio.

- 30 21A:229). In stratified epithelial tissue such as the skin, renewal occurs by mitosis of stem cells within the germinal layer, the layer closest to the basal lamina. Stem cells within the lining of the gut provide for a rapid renewal rate of this tissue. ESCs or keratinocytes obtained from the skin
- 35 or lining of the gut of a patient or donor can be grown in tissue culture (Rheinwald, 1980, Meth. Cell Bio. 21A:229; Pittelkow and Scott, 1986, Mayo Clinic Proc. 61:771). If the

ESCs are provided by a donor, a method for suppression of host versus graft reactivity (e.g., irradiation, drug or antibody administration to promote moderate immunosuppression) can also be used.

- with respect to hematopoietic stem cells (HSC), any technique which provides for the isolation, propagation, and maintenance in vitro of HSC can be used in this embodiment of the invention. Techniques by which this may be accomplished include (a) the isolation and establishment of HSC cultures
- 10 from bone marrow cells isolated from the future host, or a donor, or (b) the use of previously established long-term HSC cultures, which may be allogeneic or xenogeneic. Non-autologous HSC are used preferably in conjunction with a method of suppressing transplantation immune reactions of the
- 15 future host/patient. In a particular embodiment of the present invention, human bone marrow cells can be obtained from the posterior iliac crest by needle aspiration (see, e.g., Kodo et al., 1984, J. Clin. Invest. 73:1377-1384). In a preferred embodiment of the present invention, the HSCs can
- 20 be made highly enriched or in substantially pure form. This enrichment can be accomplished before, during, or after long-term culturing, and can be done by any techniques known in the art. Long-term cultures of bone marrow cells can be established and maintained by using, for example, modified
- 25 Dexter cell culture techniques (Dexter et al., 1977, J. Cell Physiol. 91:335) or Witlock-Witte culture techniques (Witlock and Witte, 1982, Proc. Natl. Acad. Sci. USA 79:3608-3612).

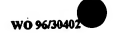
In a specific embodiment, the nucleic acid to be introduced for purposes of gene therapy comprises an 30 inducible promoter operably linked to the coding region, such that expression of the nucleic acid is controllable by controlling the presence or absence of the appropriate inducer of transcription.

Additional methods that can be adapted for use to 35 deliver a nucleic acid encoding a lats protein or functional derivative thereof are described in Section 5.8.2.2.2. WO 96/3

### 5.8.2. TREATMENT AND PREVENTION OF DISORDERS IN WHICH CELL PROLIFERATION IS DESIRED

Diseases and disorders involving a deficiency in cell proliferation (growth) or in which cell proliferation is 5 otherwise desirable for treatment or prevention, are treated or prevented by administration of a Therapeutic that antagonizes (inhibits) lats function (in particular, latsmediated inhibition of cell proliferation). Therapeutics that can be used include but are not limited to anti-lats antibodies (and fragments and derivatives thereof containing the binding region thereof), lats derivatives or analogs that are dominant-negative kinases, lats antisense nucleic acids, and lats nucleic acids that are dysfunctional (e.g., due to a heterologous (non-lats sequence) insertion within the lats 15 coding sequence) that are used to "knockout" endogenous lats function by homologous recombination (see, e.g., Capecchi, 1989, Science 244:1288-1292). In a specific embodiment of the invention, a nucleic acid containing a portion of a lats gene in which lats sequences flank (are both 5' and 3' to) a 20 different gene sequence, is used, as a lats antagonist, to promote lats inactivation by homologous recombination (see also Koller and Smithies, 1989, Proc. Natl. Acad. Sci. USA 86:8932-8935; Zijlstra et al., 1989, Nature 342:435-438). Other Therapeutics that inhibit lats function can be 25 identified by use of known convenient in vitro assays, e.g., based on their ability to inhibit binding of lats to another protein (e.g., an SH3-domain containing protein), or inhibit any known lats function, as preferably assayed in vitro or in cell culture, although genetic assays (e.g., in Drosophila) 30 may also be employed. Preferably, suitable in vitro or in vivo assays, are utilized to determine the effect of a specific Therapeutic and whether its administration is indicated for treatment of the affected tissue.

In specific embodiments, Therapeutics that inhibit lats function are administered therapeutically (including prophylactically): (1) in diseases or disorders involving an increased (relative to normal or desired) level of lats



protein or function, for example, in patients where lats protein is overactive or overexpressed; or (2) in diseases or disorders wherein in vitro (or in vivo) assays (see infra) indicate the utility of lats antagonist administration. The

- 5 increased levels in lats protein or function can be readily detected, e.g., by quantifying protein and/or RNA, by obtaining a patient tissue sample (e.g., from biopsy tissue) and assaying it in vitro for RNA or protein levels, structure and/or activity of the expressed lats RNA or protein. Many
- 10 methods standard in the art can be thus employed, including but not limited to kinase assays, immunoassays to detect and/or visualize lats protein (e.g., Western blot, immunoprecipitation followed by sodium dodecyl sulfate polyacrylamide gel electrophoresis, immunocytochemistry,
- 15 etc.) and/or hybridization assays to detect lats expression by detecting and/or visualizing respectively lats mRNA (e.g., Northern assays, dot blots, in situ hybridization, etc.), etc.

Diseases and disorders involving a deficiency in

20 cell proliferation or in which cell proliferation is desired
for treatment or prevention, and that can be treated or
prevented by inhibiting lats function, include but are not
limited to degenerative disorders, growth deficiencies,
hypoproliferative disorders, physical trauma, lesions, and
25 wounds: for example, to promote wound healing, or to promote

- 25 wounds; for example, to promote wound healing, or to promote regeneration in degenerated, lesioned or injured tissues, etc. In a specific embodiment, nervous system disorders are treated. In another specific embodiment, a disorder that is not of the nervous system is treated.
- Lesions which may be treated according to the present invention include but are not limited to the following lesions:
  - (i) traumatic lesions, including lesions caused by physical injury or associated with surgery;
- (ii) ischemic lesions, in which a lack of oxygen results in cell injury or death, e.g.,

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		myocardial or cerebral infarction or ischemia
		or spinal cord infarction or ischemia;
	(iii)	malignant lesions, in which cells are
		destroyed or injured by malignant tissue;
5	(iv)	infectious lesions, in which tissue is
		destroyed or injured as a result of infection,
		for example, by an abscess or associated with
		infection by human immunodeficiency virus,
		herpes zoster, or herpes simplex virus or with
10		Lyme disease, tuberculosis, syphilis;
	(v)	degenerative lesions, in which tissue is
		destroyed or injured as a result of a
		degenerative process, including but not
		limited to nervous system degeneration
15		associated with Parkinson's disease,
		Alzheimer's disease, Huntington's chorea, or
		amyotrophic lateral sclerosis;
	(vi)	lesions associated with nutritional diseases
		or disorders, in which tissue is destroyed or
20		injured by a nutritional disorder or disorder
		of metabolism including but not limited to,
		vitamin B12 deficiency, folic acid deficiency,
		Wernicke disease, tobacco-alcohol amblyopia,
		Marchiafava-Bignami disease (primary
25		degeneration of the corpus callosum), and
		alcoholic cerebellar degeneration;
	(vii)	lesions associated with systemic diseases
		including but not limited to diabetes or
		systemic lupus erythematosus;
30	(viii)	lesions caused by toxic substances including
		alcohol, lead, or other toxins; and
	(ix)	demyelinated lesions of the nervous system, in
		which a portion of the nervous system is
		destroyed or injured by a demyelinating
15		disease including but not limited to multiple
		sclerosis, human immunodeficiency virus-
		associated myelopathy, transverse myelopathy

or various etiologies, progressive multifocal leukoencephalopathy, and central pontine myelinolysis.

Nervous system lesions which may be treated in a 5 patient (including human and non-human mammalian patients) according to the invention include but are not limited to the lesions of either the central (including spinal cord, brain) or peripheral nervous systems.

Therapeutics which are useful according to this

10 embodiment of the invention for treatment of a disorder may
be selected by testing for biological activity in promoting
the survival or differentiation of cells (see also Section
5.9). For example, in a specific embodiment relating to
therapy of the nervous system, a Therapeutic which elicits

15 one of the following effects may be useful according to the
invention:

- (i) increased sprouting of neurons in culture or in vivo;
- (ii) increased production of a neuron-associated

  molecule in culture or in vivo, e.g., choline
  acetyltransferase or acetylcholinesterase with
  respect to motor neurons; or
  - (iii) decreased symptoms of neuron dysfunction in vivo.
- In preferred, non-limiting embodiments, increased sprouting of neurons may be detected by methods set forth in Pestronk et al. (1980, Exp. Neurol. 70:65-82) or Brown et al. (1981, Ann. Rev. Neurosci. 4:17-42); and increased production of neuron-associated molecules may be measured by bioassay, enzymatic assay, antibody binding, Northern blot assay, etc.,
- 5.8.2.1. ANTISENSE REGULATION OF LATS EXPRESSION

  In a specific embodiment, lats function is
  inhibited by use of lats antisense nucleic acids. The

depending on the molecule to be measured.

present invention provides the therapeutic or prophylactic use of nucleic acids of at least six nucleotides that are antisense to a gene or cDNA encoding lats or a portion thereof. A lats "antisense" nucleic acid as used herein

- 5 refers to a nucleic acid capable of hybridizing to a portion of a lats RNA (preferably mRNA) by virtue of some sequence complementarity. The antisense nucleic acid may be complementary to a coding and/or noncoding region of a lats mRNA. Such antisense nucleic acids have utility as
- 10 Therapeutics that inhibits lats function, and can be used in the treatment or prevention of disorders as described *supra* in Section 5.8.2 and its subsections.

The antisense nucleic acids of the invention can be oligonucleotides that are double-stranded or single-stranded,

15 RNA or DNA or a modification or derivative thereof, which can be directly administered to a cell, or which can be produced intracellularly by transcription of exogenous, introduced sequences.

In a specific embodiment, the lats antisense

20 nucleic acids provided by the instant invention can be used
to promote regeneration or wound healing or to promote growth
(larger size).

The invention further provides pharmaceutical compositions comprising an effective amount of the lats

25 antisense nucleic acids of the invention in a pharmaceutically acceptable carrier, as described infra.

In another embodiment, the invention is directed to methods for inhibiting the expression of a lats nucleic acid sequence in a prokaryotic or eukaryotic cell comprising

30 providing the cell with an effective amount of a composition comprising an lats antisense nucleic acid of the invention.

Lats antisense nucleic acids and their uses are described in detail below.

### 5.8.2.1.1. LATS ANTISENSE NUCLEIC ACIDS

The lats antisense nucleic acids are of at least six nucleotides and are preferably oligonucleotides (ranging from 6 to about 50 oligonucleotides). In specific aspects,

- 5 the oligonucleotide is at least 10 nucleotides, at least 15 nucleotides, at least 100 nucleotides, or at least 200 nucleotides. The oligonucleotides can be DNA or RNA or chimeric mixtures or derivatives or modified versions thereof, single-stranded or double-stranded. The
- noiety, or phosphate backbone. The oligonucleotide may include other appending groups such as peptides, or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc. Natl. Acad. Sci. U.S.A.
- 15 86:6553-6556; Lemaitre et al., 1987, Proc. Natl. Acad. Sci. 84:648-652; PCT Publication No. WO 88/09810, published December 15, 1988) or blood-brain barrier (see, e.g., PCT Publication No. WO 89/10134, published April 25, 1988), hybridization-triggered cleavage agents (see, e.g., Krol et
- 20 al., 1988, BioTechniques 6:958-976) or intercalating agents (see, e.g., Zon, 1988, Pharm. Res. 5:539-549).

In a preferred aspect of the invention, a lats antisense oligonucleotide is provided, preferably of single-stranded DNA. In a most preferred aspect, such an

- 25 oligonucleotide comprises a sequence antisense to the sequence encoding an SH3 binding domain or a kinase domain of a lats protein, most preferably, of a human lats protein. The oligonucleotide may be modified at any position on its structure with substituents generally known in the art.
- least one modified base moiety which is selected from the group including but not limited to 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xantine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil,
- 35 5-carboxymethylaminomethyl-2-thiouridine,
  5-carboxymethylaminomethyluracil, dihydrouracil, beta-Dgalactosylqueosine, inosine, N6-isopentenyladenine,



1-methylguanine, 1-methylinosine, 2,2-dimethylguanine,
2-methyladenine, 2-methylguanine, 3-methylcytosine,
5-methylcytosine, N6-adenine, 7-methylguanine,
5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil,
5 beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil,
5-methoxyuracil, 2-methylthio-N6-isopentenyladenine,
uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil,
queosine, 2-thiocytosine, 5-methyl-2-thiouracil,
2-thiouracil, 4-thiouracil, 5-methyluracil, uracil10 5-oxyacetic acid methylester, uracil-5-oxyacetic acid (v),
5-methyl-2-thiouracil, 3-(3-amino-3-N-2-carboxypropyl)

uracil, (acp3)w, and 2,6-diaminopurine.

In another embodiment, the oligonucleotide comprises at least one modified sugar moiety selected from

15 the group including but not limited to arabinose, 2-fluoroarabinose, xylulose, and hexose.

In yet another embodiment, the oligonucleotide comprises at least one modified phosphate backbone selected from the group consisting of a phosphorothicate, a

20 phosphorodithioate, a phosphoramidothioate, a phosphoramidate, a phosphordiamidate, a methylphosphonate, an alkyl phosphotriester, and a formacetal or analog thereof.

In yet another embodiment, the oligonucleotide is an  $\alpha$ -anomeric oligonucleotide. An  $\alpha$ -anomeric oligonucleotide **25** forms specific double-stranded hybrids with complementary RNA in which, contrary to the usual  $\beta$ -units, the strands run parallel to each other (Gautier et al., 1987, Nucl. Acids Res. 15:6625-6641).

The oligonucleotide may be conjugated to another 30 molecule, e.g., a peptide, hybridization triggered cross-linking agent, transport agent, hybridization-triggered cleavage agent, etc.

Oligonucleotides of the invention may be synthesized by standard methods known in the art, e.g. by use 35 of an automated DNA synthesizer (such as are commercially available from Biosearch, Applied Biosystems, etc.). As examples, phosphorothicate oligonucleotides may be

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synthesized by the method of Stein et al. (1988, Nucl. Acids Res. 16:3209), methylphosphonate oligonucleotides can be prepared by use of controlled pore glass polymer supports (Sarin et al., 1988, Proc. Natl. Acad. Sci. U.S.A. 85:7448-57451), etc.

In a specific embodiment, the *lats* antisense oligonucleotide comprises catalytic RNA, or a ribozyme (see, e.g., PCT International Publication WO 90/11364, published October 4, 1990; Sarver et al., 1990, Science 247:1222-1225).

10 In another embodiment, the oligonucleotide is a 2'-0-methylribonucleotide (Inoue et al., 1987, Nucl. Acids Res. 15:6131-6148), or a chimeric RNA-DNA analogue (Inoue et al., 1987, FEBS Lett. 215:327-330).

In an alternative embodiment, the lats antisense

15 nucleic acid of the invention is produced intracellularly by
transcription from an exogenous sequence. For example, a
vector can be introduced in vivo such that it is taken up by
a cell, within which cell the vector or a portion thereof is
transcribed, producing an antisense nucleic acid (RNA) of the

- 20 invention. Such a vector would contain a sequence encoding the lats antisense nucleic acid. Such a vector can remain episomal or become chromosomally integrated, as long as it can be transcribed to produce the desired antisense RNA. Such vectors can be constructed by recombinant DNA technology
- 25 methods standard in the art. Vectors can be plasmid, viral, or others known in the art, used for replication and expression in mammalian cells. Expression of the sequence encoding the lats antisense RNA can be by any promoter known in the art to act in mammalian, preferably human, cells.
- 30 Such promoters can be inducible or constitutive. Such promoters include but are not limited to: the SV40 early promoter region (Bernoist and Chambon, 1981, Nature 290:304-310), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto et al., 1980, Cell 22:787-
- 35 797), the herpes thymidine kinase promoter (Wagner et al., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:1441-1445), the



regulatory sequences of the metallothionein gene (Brinster et al., 1982, Nature 296:39-42), etc.

The antisense nucleic acids of the invention comprise a sequence complementary to at least a portion of an 5 RNA transcript of a lats gene, preferably a human lats gene. However, absolute complementarity, although preferred, is not required. A sequence "complementary to at least a portion of an RNA," as referred to herein, means a sequence having sufficient complementarity to be able to hybridize with the 10 RNA, forming a stable duplex; in the case of double-stranded lats antisense nucleic acids, a single strand of the duplex DNA may thus be tested, or triplex formation may be assayed. The ability to hybridize will depend on both the degree of complementarity and the length of the antisense nucleic acid. 15 Generally, the longer the hybridizing nucleic acid, the more base mismatches with a lats RNA it may contain and still form a stable duplex (or triplex, as the case may be). One skilled in the art can ascertain a tolerable degree of mismatch by use of standard procedures to determine the 20 melting point of the hybridized complex.

### 5.8.2.1.2. THERAPEUTIC USE OF LATS ANTISENSE NUCLEIC ACIDS

The lats antisense nucleic acids can be used to

treat (or prevent) disorders of a cell type that expresses,
or preferably overexpresses, lats. In a specific embodiment,
such a disorder is a growth deficiency. In a preferred
embodiment, a single-stranded DNA antisense lats
oligonucleotide is used.

can be identified by various methods known in the art. Such methods include but are not limited to hybridization with a lats-specific nucleic acid (e.g. by Northern hybridization, dot blot hybridization, in situ hybridization), observing the ability of RNA from the cell type to be translated in vitro into lats, immunoassay, etc. In a preferred aspect, primary tissue from a patient can be assayed for lats expression

prior to treatment, e.g., by immunocytochemistry or in situ hybridization.

Pharmaceutical compositions of the invention (see Section 5.10), comprising an effective amount of a lats

santisense nucleic acid in a pharmaceutically acceptable carrier, can be administered to a patient having a disease or disorder which is of a type that expresses or overexpresses lats RNA or protein.

The amount of lats antisense nucleic acid which

10 will be effective in the treatment of a particular disorder
or condition will depend on the nature of the disorder or
condition, and can be determined by standard clinical
techniques. Where possible, it is desirable to determine the
antisense cytotoxicity of the tumor type to be treated in

15 vitro, and then in useful animal model systems prior to
testing and use in humans.

In a specific embodiment, pharmaceutical compositions comprising lats antisense nucleic acids are administered via liposomes, microparticles, or microcapsules.

- 20 In various embodiments of the invention, it may be useful to use such compositions to achieve sustained release of the lats antisense nucleic acids. In a specific embodiment, it may be desirable to utilize liposomes targeted via antibodies to specific identifiable tumor antigens (Leonetti et al.,
- 25 1990, Proc. Natl. Acad. Sci. U.S.A. 87:2448-2451; Renneisen et al., 1990, J. Biol. Chem. 265:16337-16342).

Additional methods that can be adapted for use to deliver a lats antisense nucleic acid are described in Section 5.8.1.4.

30

## 5.9. DEMONSTRATION OF THERAPEUTIC OR PROPHYLACTIC UTILITY

The Therapeutics of the invention are preferably tested in vitro, and then in vivo for the desired therapeutic or prophylactic activity, prior to use in humans.

For example, In vitro assays which can be used to determine whether administration of a specific Therapeutic is



indicated, include in vitro cell culture assays in which a patient tissue sample is grown in culture, and exposed to or otherwise administered a Therapeutic, and the effect of such Therapeutic upon the tissue sample is observed. In one

- 5 embodiment, where the patient has a malignancy, a sample of cells from such malignancy is plated out or grown in culture, and the cells are then exposed to a Therapeutic. A Therapeutic which inhibits survival or growth of the malignant cells is selected for therapeutic use in vivo.
- 10 Many assays standard in the art can be used to assess such survival and/or growth; for example, cell proliferation can be assayed by measuring <sup>3</sup>H-thymidine incorporation, by direct cell count, by detecting changes in transcriptional activity of known genes such as proto-oncogenes (e.g., fos, myc) or
- 15 cell cycle markers; cell viability can be assessed by trypan blue staining, differentiation can be assessed visually based on changes in morphology, etc.

In another embodiment, a Therapeutic is indicated for use which exhibits the desired effect, inhibition or 20 promotion of cell growth, upon a patient cell sample from tissue having or suspected of having a hyper- or hypoproliferative disorder, respectively. Such hyper- or hypoproliferative disorders include but are not limited to those described in Sections 5.8.1 through 5.8.3 infra.

In another specific embodiment, a Therapeutic is indicated for use in treating cell injury or a degenerative disorder (see Section 5.8.2) which exhibits in vitro promotion of growth/proliferation of cells of the affected patient type. Regarding nervous system disorders, see also Section 5.8.2.1 for assays that can be used.

In various specific embodiments, in vitro assays can be carried out with representative cells of cell types involved in a patient's disorder, to determine if a Therapeutic has a desired effect upon such cell types.

In another embodiment, cells of a patient tissue sample suspected of being pre-neoplastic are similarly plated out or grown in vitro, and exposed to a Therapeutic. The

Therapeutic which results in a cell phenotype that is more normal (i.e., less representative of a pre-neoplastic state, neoplastic state, malignant state, or transformed phenotype) is selected for therapeutic use. Many assays standard in the 5 art can be used to assess whether a pre-neoplastic state, neoplastic state, or a transformed or malignant phenotype, is present. For example, characteristics associated with a transformed phenotype (a set of in vitro characteristics associated with a tumorigenic ability in vivo) include a more 10 rounded cell morphology, looser substratum attachment, loss of contact inhibition, loss of anchorage dependence, release of proteases such as plasminogen activator, increased sugar transport, decreased serum requirement, expression of fetal antigens, disappearance of the 250,000 dalton surface 15 protein, etc. (see Luria et al., 1978, General Virology, 3d Ed., John Wiley & Sons, New York pp. 436-446).

In other specific embodiments, the in vitro assays described supra can be carried out using a cell line, rather than a cell sample derived from the specific patient to be treated, in which the cell line is derived from or displays characteristic(s) associated with the malignant, neoplastic or pre-neoplastic disorder desired to be treated or prevented, or is derived from the cell type upon which an effect is desired, according to the present invention.

suitable animal model systems prior to testing in humans, including but not limited to rats, mice, chicken, cows, monkeys, rabbits, etc. For in vivo testing, prior to administration to humans, any animal model system known in the art may be used.

### 5.10. THERAPEUTIC/PROPHYLACTIC ADMINISTRATION AND COMPOSITIONS

The invention provides methods of treatment (and prophylaxis) by administration to a subject of an effective amount of a Therapeutic of the invention. In a preferred aspect, the Therapeutic is substantially purified. The

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subject is preferably an animal, including but not limited to animals such as cows, pigs, horses, chickens, cats, dogs, etc., and is preferably a mammal, and most preferably human. In a specific embodiment, a non-human mammal is the subject.

- Formulations and methods of administration that can be employed when the Therapeutic comprises a nucleic acid are described in Sections 5.8.1.4 and 5.8.2.2 above; additional appropriate formulations and routes of administration can be selected from among those described hereinbelow.
- Various delivery systems are known and can be used to administer a Therapeutic of the invention, e.g., encapsulation in liposomes, microparticles, microcapsules, recombinant cells capable of expressing the Therapeutic, receptor-mediated endocytosis (see, e.g., Wu and Wu, 1987, J.
- 15 Biol. Chem. 262:4429-4432), construction of a Therapeutic nucleic acid as part of a retroviral or other vector, etc. Methods of introduction include but are not limited to intradermal, intramuscular, intraperitoneal, intravenous, subcutaneous, intranasal, epidural, and oral routes. The
- 20 compounds may be administered by any convenient route, for example by infusion or bolus injection, by absorption through epithelial or mucocutaneous linings (e.g., oral mucosa, rectal and intestinal mucosa, etc.) and may be administered together with other biologically active agents.
- 25 Administration can be systemic or local. In addition, it may be desirable to introduce the pharmaceutical compositions of the invention into the central nervous system by any suitable route, including intraventricular and intrathecal injection; intraventricular injection may be facilitated by an
- 30 intraventricular catheter, for example, attached to a reservoir, such as an Ommaya reservoir. Pulmonary administration can also be employed, e.g., by use of an inhaler or nebulizer, and formulation with an aerosolizing agent.
- In a specific embodiment, it may be desirable to administer the pharmaceutical compositions of the invention locally to the area in need of treatment; this may be

achieved by, for example, and not by way of limitation, local infusion during surgery, topical application, e.g., in conjunction with a wound dressing after surgery, by injection, by means of a catheter, by means of a suppository, 5 or by means of an implant, said implant being of a porous, non-porous, or gelatinous material, including membranes, such as sialastic membranes, or fibers. In one embodiment, administration can be by direct injection at the site (or former site) of a malignant tumor or neoplastic or pre-

10 neoplastic tissue. In another embodiment, the Therapeutic can be delivered in a vesicle, in particular a liposome (see Langer, Science 249:1527-1533 (1990); Treat et al., in Liposomes in the Therapy of Infectious Disease and Cancer, Lopez-Berestein 15 and Fidler (eds.), Liss, New York, pp. 353-365 (1989); Lopez-Berestein, ibid., pp. 317-327; see generally ibid.)

In yet another embodiment, the Therapeutic can be delivered in a controlled release system. In one embodiment, a pump may be used (see Langer, supra; Sefton, CRC Crit. Ref.

- 20 Biomed. Eng. 14:201 (1987); Buchwald et al., Surgery 88:507 (1980); Saudek et al., N. Engl. J. Med. 321:574 (1989)). In another embodiment, polymeric materials can be used (see Medical Applications of Controlled Release, Langer and Wise (eds.), CRC Pres., Boca Raton, Florida (1974); Controlled
- 25 Drug Bioavailability, Drug Product Design and Performance, Smolen and Ball (eds.), Wiley, New York (1984); Ranger and Peppas, J. Macromol. Sci. Rev. Macromol. Chem. 23:61 (1983); see also Levy et al., Science 228:190 (1985); During et al., Ann. Neurol. 25:351 (1989); Howard et al., J. Neurosurg.
- 30 71:105 (1989)). In yet another embodiment, a controlled release system can be placed in proximity of the therapeutic target, i.e., the brain, thus requiring only a fraction of the systemic dose (see, e.g., Goodson, in Medical Applications of Controlled Release, supra, vol. 2, pp.
- **35** 115-138 (1984)).

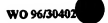
Other controlled release systems are discussed in the review by Langer (Science 249:1527-1533 (1990)).

homologous recombination.

In a specific embodiment where the Therapeutic is a nucleic acid encoding a protein Therapeutic, the nucleic acid can be administered in vivo to promote expression of its encoded protein, by constructing it as part of an appropriate nucleic acid expression vector and administering it so that it becomes intracellular, e.g., by use of a retroviral vector (see U.S. Patent No. 4,980,286), or by direct injection, or by use of microparticle bombardment (e.g., a gene gun; Biolistic, Dupont), or coating with lipids or cell-surface receptors or transfecting agents, or by administering it in linkage to a homeobox-like peptide which is known to enter the nucleus (see e.g., Joliot et al., 1991, Proc. Natl. Acad. Sci. USA 88:1864-1868), etc. Alternatively, a nucleic acid Therapeutic can be introduced intracellularly and incorporated within host cell DNA for expression, by

The present invention also provides pharmaceutical compositions. Such compositions comprise a therapeutically effective amount of a Therapeutic, and a pharmaceutically

- 20 acceptable carrier. In a specific embodiment, the term "pharmaceutically acceptable" means approved by a regulatory agency of the Federal or a state government or listed in the U.S. Pharmacopeia or other generally recognized pharmacopeia for use in animals, and more particularly in humans. The
- 25 term "carrier" refers to a diluent, adjuvant, excipient, or vehicle with which the therapeutic is administered. Such pharmaceutical carriers can be sterile liquids, such as water and oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral
- 30 oil, sesame oil and the like. Water is a preferred carrier when the pharmaceutical composition is administered intravenously. Saline solutions and aqueous dextrose and glycerol solutions can also be employed as liquid carriers, particularly for injectable solutions. Suitable
- 35 pharmaceutical excipients include starch, glucose, lactose, sucrose, gelatin, malt, rice, flour, chalk, silica gel, sodium stearate, glycerol monostearate, talc, sodium



chloride, dried skim milk, glycerol, propylene, glycol, water, ethanol and the like. The composition, if desired, can also contain minor amounts of wetting or emulsifying agents, or pH buffering agents. These compositions can take

- 5 the form of solutions, suspensions, emulsion, tablets, pills, capsules, powders, sustained-release formulations and the like. The composition can be formulated as a suppository, with traditional binders and carriers such as triglycerides. Oral formulation can include standard carriers such as
- 10 pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate, etc. Examples of suitable pharmaceutical carriers are described in "Remington's Pharmaceutical Sciences" by E.W. Martin. Such compositions will contain a therapeutically
- 15 effective amount of the Therapeutic, preferably in purified form, together with a suitable amount of carrier so as to provide the form for proper administration to the patient.

  The formulation should suit the mode of administration.

In a preferred embodiment, the composition is

- 20 formulated in accordance with routine procedures as a pharmaceutical composition adapted for intravenous administration to human beings. Typically, compositions for intravenous administration are solutions in sterile isotonic aqueous buffer. Where necessary, the composition may also
- 25 include a solubilizing agent and a local anesthetic such as lignocaine to ease pain at the site of the injection. Generally, the ingredients are supplied either separately or mixed together in unit dosage form, for example, as a dry lyophilized powder or water free concentrate in a
- 30 hermetically sealed container such as an ampoule or sachette indicating the quantity of active agent. Where the composition is to be administered by infusion, it can be dispensed with an infusion bottle containing sterile pharmaceutical grade water or saline. Where the composition
- 35 is administered by injection, an ampoule of sterile water for injection or saline can be provided so that the ingredients may be mixed prior to administration.



The Therapeutics of the invention can be formulated as neutral or salt forms. Pharmaceutically acceptable salts include those formed with free amino groups such as those derived from hydrochloric, phosphoric, acetic, oxalic,

- 5 tartaric acids, etc., and those formed with free carboxyl groups such as those derived from sodium, potassium, ammonium, calcium, ferric hydroxides, isopropylamine, triethylamine, 2-ethylamino ethanol, histidine, procaine, etc.
- The amount of the Therapeutic of the invention which will be effective in the treatment of a particular disorder or condition will depend on the nature of the disorder or condition, and can be determined by standard clinical techniques. In addition, in vitro assays may
- 15 optionally be employed to help identify optimal dosage ranges. The precise dose to be employed in the formulation will also depend on the route of administration, and the seriousness of the disease or disorder, and should be decided according to the judgment of the practitioner and each
- 20 patient's circumstances. However, suitable dosage ranges for intravenous administration are generally about 20-500 micrograms of active compound per kilogram body weight. Suitable dosage ranges for intranasal administration are generally about 0.01 pg/kg body weight to 1 mg/kg body
- 25 weight. Effective doses may be extrapolated from doseresponse curves derived from in vitro or animal model test systems.

Suppositories generally contain active ingredient in the range of 0.5% to 10% by weight; oral formulations 30 preferably contain 10% to 95% active ingredient.

The invention also provides a pharmaceutical pack or kit comprising one or more containers filled with one or more of the ingredients of the pharmaceutical compositions of the invention. Optionally associated with such container(s)

35 can be a notice in the form prescribed by a governmental agency regulating the manufacture, use or sale of pharmaceuticals or biological products, which notice reflects



approval by the agency of manufacture, use or sale for human administration.

#### 5.11. ADDITIONAL USE OF INHIBITION OF LATS FUNCTION TO PROMOTE INCREASED GROWTH

5 Inhibition of lats function (e.g., by administering a compound that inhibits lats function as described in Sections 5.8.2 through 5.8.2.1.2 above), has utility that is not limited to therapeutic or prophylactic applications. example, lats function can be inhibited in order to increase growth of animals (e.g., cows, horses, pigs, goats, deer, chickens) and plants (particularly edible plants, e.g., tomatoes, melons, lettuce, carrots, potatoes, and other vegetables), particularly those that are food or material sources. For example, antisense inhibition (preferably where the lats antisense nucleic acid is under the control of a tissue-specific promoter) can be used in plants or animals to increase growth where desired (e.g., in the fruit or muscle). For example, a lats antisense nucleic acid under the control of a temperature-sensitive promoter can be administered to a plant or animal, and the desired portion of the (or the entire) plant or animal can be subjected to heat in order to induce antisense nucleic acid production, resulting lats inhibition, and resulting cell proliferation. embodiments, chemical mutagenesis, or homologous recombination with an insertionally inactivated lats gene (see Capecchi, 1989, Science 244:1288-1292 and Section 5.14 infra) can be carried out to reduce or destroy endogenous lats function, in order to achieve increased growth. Suitable methods, modes of administration and compositions, that can be used to inhibit lats function are described in Sections 5.8.2 through 5.8.2.1.2, above. Methods to make plants recombinant are commonly known in the art and can be Regarding methods of plant transformation (e.g., for 35 transformation with a lats antisense nucleic acid or with a sequence encoding a lats derivative that is a dominant-

negative kinase), see e.g., Valvekens et al., 1988, Proc.

Natl. Acad. Sci. USA 85:5536-5540. Regarding methods of targeted gene inactivation in plants (e.g., to inactivate lats), see e.g., Miao and Lam, 1995, The Plant J. 7:359-365.

Inhibition of lats function can also have uses in vitro, e.g., to expand cells in vitro, including but not limited to stem cells, progenitor cells, muscle cells, fibroblasts, liver cells, etc., e.g., to grow cells/tissue in vitro prior to administration to a patient (preferably a patient from which the cells were derived), etc.

10

## 5.12. ADDITIONAL USE OF INHIBITION OF LATS FUNCTION TO INHIBIT CELLULAR SENESCENCE

Inhibition of lats function (e.g., by administering a compound that inhibits lats function as described in Sections 5.8.2 through 5.8.2.1.2 above), also has utility in the inhibition of cellular senescence. Thus, inhibition of lats function can be carried out to delay or prevent the onset of cellular senescence, in vivo or in vitro. In a specific embodiment, cellular senescence is delayed or prevented without incurring the onset of cell malignancy or its in vitro correlate, a transformed phenotype.

Thus, for example, a lats antagonist (e.g., antilats antibody, lats derivatives or analogs that are dominantnegative kinases; lats antisense nucleic acids, etc.) can be
administered to a subject to inhibit or prevent aging or cell
death or the effects of aging or cell death (e.g., in the
skin, wrinkling, loss of elasticity, less uniform skin tone;
in the skin and elsewhere, loss of known characteristics of
proper physiological function such as expression of
characteristic antigens, secreted molecules, etc.) In one
embodiment, a lats antagonist is applied topically, e.g., in
a cream or gel, to the skin of the subject. In another
embodiment, a lats antagonist is injected, e.g.,
intradermally, intraperitoneally, or intramuscularly.

In a specific embodiment, a lats antagonist is contacted with cells grown in culture, e.g., by addition of the antagonist to the culture medium or by adsorption of the



antagonist to the culture plate or flask prior to seeding of the cells, in order to inhibit or delay senescence in vitro, e.g., to delay "crisis" phase. For example, such a method can be carried out in order to lengthen the time that cells

- 5 can be kept alive in vitro, e.g., in order to facilitate conducting studies of the toxicity of a compound (e.g., a lead drug candidate) upon such cells, to study the effect of a molecule upon cell function, and, generally, to study the function of such cells. Such cells include but are not
- 10 limited to neurons of the central nervous system (e.g., hippocampal, hypothalmic) or peripheral nervous system, glial cells, fibroblasts, kidney cells, liver cells, heart cells, muscle cells, endothelial cells, melanocytes, and hematopoietic cells such and T and B lymphocytes,

In vitro assays of senescence are well known in the art and can be used to screen potential lats antagonists prior to use in this aspect of the invention (see, e.g., Hubbard and Ozer, 1995, "Senescence and immortalization of

20 human cells," in <u>Cell Growth and Apoptosis</u>. A <u>Practical Approach</u>, Ch. 12, Studzinski, G.P. (ed.), Oxford University Press, Inc., New York, NY, pp. 229-248.

#### 5.13. **DIAGNOSIS AND SCREENING**

subsequences thereof, lats nucleic acids (and sequences complementary thereto), anti-lats antibodies, have uses in diagnostics. Such molecules can be used in assays, such as immunoassays, to detect, prognose, diagnose, or monitor various conditions, diseases, and disorders affecting lats expression, or monitor the treatment thereof. In particular, such an immunoassay is carried out by a method comprising contacting a sample derived from a patient with an anti-lats antibody under conditions such that immunospecific binding can occur, and detecting or measuring the amount of any immunospecific binding by the antibody. In a specific

aspect, such binding of antibody, in tissue sections, can be



used to detect aberrant lats localization or aberrant (e.g., low or absent) levels of lats. In a specific embodiment, antibody to lats can be used to assay in a patient tissue or serum sample for the presence of lats where an aberrant level

- 5 of lats is an indication of a diseased condition. By "aberrant levels," is meant increased or decreased levels relative to that present, or a standard level representing that present, in an analogous sample from a portion of the body or from a subject not having the disorder.
- The immunoassays which can be used include but are not limited to competitive and non-competitive assay systems using techniques such as western blots, radioimmunoassays, ELISA (enzyme linked immunosorbent assay), "sandwich" immunoassays, immunoprecipitation assays, precipitin
- 15 reactions, gel diffusion precipitin reactions, immunodiffusion assays, agglutination assays, complementfixation assays, immunoradiometric assays, fluorescent immunoassays, protein A immunoassays, to name but a few.

Lats genes and related nucleic acid sequences and subsequences, including complementary sequences, can also be used in hybridization assays. Lats nucleic acid sequences, or subsequences thereof comprising about at least 8 nucleotides, can be used as hybridization probes. Hybridization assays can be used to detect, prognose,

- 25 diagnose, or monitor conditions, disorders, or disease states associated with aberrant changes in lats expression and/or activity as described supra. In particular, such a hybridization assay is carried out by a method comprising contacting a sample containing nucleic acid with a nucleic
- 30 acid probe capable of hybridizing to lats DNA or RNA, under conditions such that hybridization can occur, and detecting or measuring any resulting hybridization.

In specific embodiments, diseases and disorders involving overproliferation of cells can be diagnosed, or 35 their suspected presence can be screened for, or a predisposition to develop such disorders can be detected, by detecting decreased levels of lats protein, lats RNA, or lats

functional activity (e.g., kinase activity, SH3 domainbinding activity, etc.), or by detecting mutations in lats
RNA, DNA or protein (e.g., translocations in lats nucleic
acids, truncations in the lats gene or protein, changes in
nucleotide or amino acid sequence relative to wild-type lats)
that cause decreased expression or activity of lats. Such
diseases and disorders include but are not limited to those
described in Section 5.8.1 and its subsections. By way of
example, levels of lats protein can be detected by
hybridization assays (e.g., Northern blots, dot blots), lats
kinase activity can be measured by kinase assays commonly
known in the art, lats binding to an SH3 domain-containing
protein can be done by binding assays commonly known in the

- 15 art, translocations and point mutations in lats nucleic acids can be detected by Southern blotting, RFLP analysis, PCR using primers that preferably generate a fragment spanning at least most of the lats gene, sequencing of the lats genomic DNA or cDNA obtained from the patient, etc.
- 20 In a preferred embodiment, levels of lats mRNA or protein in a patient sample are detected or measured, in which decreased levels indicate that the subject has, or has a predisposition to developing, a malignancy or hyperproliferative disorder; in which the decreased levels are relative to the levels present in an analogous sample from a portion of the body or from a subject not having the malignancy or hyperproliferative disorder, as the case may be.

In another specific embodiment, diseases and

disorders involving a deficiency in cell proliferation or in which cell proliferation is desirable for treatment, are diagnosed, or their suspected presence can be screened for, or a predisposition to develop such disorders can be detected, by detecting increased levels of lats protein, lats RNA, or lats functional activity (e.g., kinase activity, SH3 domain binding activity, etc.), or by detecting mutations in lats RNA, DNA or protein (e.g., translocations in lats

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nucleic acids, truncations in the gene or protein, changes in nucleotide or amino acid sequence relative to wild-type lats) that cause increased expression or activity of lats. Such diseases and disorders include but are not limited to those described in Section 5.8.2 and its subsections. By way of example, levels of lats protein, levels of lats RNA, lats kinase activity, lats binding activity, and the presence of translocations or point mutations can be determined as described above.

- In a specific embodiment, levels of lats mRNA or protein in a patient sample are detected or measured, in which increased levels indicate that the subject has, or has a predisposition to developing, a growth deficiency or degenerative or hypoproliferative disorder; in which the increased levels are relative to the levels present in an analogous sample from a portion of the body or from a subject not having the growth deficiency, degenerative, or hypoproliferative disorder, as the case may be.
- Kits for diagnostic use are also provided, that 20 comprise in one or more containers an anti-lats antibody, and, optionally, a labeled binding partner to the antibody. Alternatively, the anti-lats antibody can be labeled (with a detectable marker, e.g., a chemiluminescent, enzymatic, fluorescent, or radioactive moiety). A kit is also provided 25 that comprises in one or more containers a nucleic acid probe capable of hybridizing to lats RNA. In a specific embodiment, a kit can comprise in one or more containers a pair of primers (e.g., each in the size range of 6-30nucleotides) that are capable of priming amplification [e.g., 30 by polymerase chain reaction (see e.g., Innis et al., 1990, PCR Protocols, Academic Press, Inc., San Diego, CA), ligase chain reaction (see EP 320,308) use of  $Q\beta$  replicase, cyclic probe reaction, or other methods known in the art] under appropriate reaction conditions of at least a portion of a 35 lats nucleic acid. A kit can optionally further comprise in a container a predetermined amount of a purified lats protein or nucleic acid, e.g., for use as a standard or control.

5.14. SCREENING FOR LATS AGONISTS AND ANTAGONISTS

Lats nucleic acids, proteins, and derivatives also
have uses in screening assays to detect molecules that
specifically bind to lats nucleic acids, proteins, or
5 derivatives and thus have potential use as agonists or
antagonists of lats, in particular, molecules that thus

antagonists of lats, in particular, molecules that thus affect cell proliferation. In a preferred embodiment, such assays are performed to screen for molecules with potential utility as anti-cancer drugs or lead compounds for drug

10 development. The invention thus provides assays to detect molecules that specifically bind to lats nucleic acids, proteins, or derivatives. For example, recombinant cells expressing lats nucleic acids can be used to recombinantly produce lats proteins in these assays, to screen for

15 molecules that bind to a lats protein. Molecules (e.g., putative binding partners of lats) are contacted with the lats protein (or fragment thereof) under conditions conducive to binding, and then molecules that specifically bind to the lats protein are identified. Similar methods can be used to

20 screen for molecules that bind to lats derivatives or nucleic acids. Methods that can be used to carry out the foregoing are commonly known in the art.

By way of example, diversity libraries, such as random or combinatorial peptide or nonpeptide libraries can 25 be screened for molecules that specifically bind to lats.

Many libraries are known in the art that can be used, e.g., chemically synthesized libraries, recombinant (e.g., phage display libraries), and in vitro translation-based libraries.

Examples of chemically synthesized libraries are

30 described in Fodor et al., 1991, Science 251:767-773;

Houghten et al., 1991, Nature 354:84-86; Lam et al., 1991,

Nature 354:82-84; Medynski, 1994, Bio/Technology 12:709-710;

Gallop et al., 1994, J. Medicinal Chemistry 37(9):1233-1251;

Ohlmeyer et al., 1993, Proc. Natl. Acad. Sci. USA

35 90:10922-10926; Erb et al., 1994, Proc. Natl. Acad. Sci. USA 91:11422-11426; Houghten et al., 1992, Biotechniques 13:412; Jayawickreme et al., 1994, Proc. Natl. Acad. Sci. USA



91:1614-1618; Salmon et al., 1993, Proc. Natl. Acad. Sci. USA 90:11708-11712; PCT Publication No. WO 93/20242; and Brenner and Lerner, 1992, Proc. Natl. Acad. Sci. USA 89:5381-5383.

Examples of phage display libraries are described 5 in Scott and Smith, 1990, Science 249:386-390; Devlin et al., 1990, Science, 249:404-406; Christian, R.B., et al., 1992, J. Mol. Biol. 227:711-718); Lenstra, 1992, J. Immunol. Meth. 152:149-157; Kay et al., 1993, Gene 128:59-65; and PCT Publication No. WO 94/18318 dated August 18, 1994.

In vitro translation-based libraries include but are not limited to those described in PCT Publication No. WO 91/05058 dated April 18, 1991; and Mattheakis et al., 1994, Proc. Natl. Acad. Sci. USA 91:9022-9026.

By way of examples of nonpeptide libraries, a

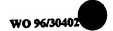
15 benzodiazepine library (see e.g., Bunin et al., 1994, Proc.
Natl. Acad. Sci. USA 91:4708-4712) can be adapted for use.
Peptoid libraries (Simon et al., 1992, Proc. Natl. Acad. Sci.
USA 89:9367-9371) can also be used. Another example of a
library that can be used, in which the amide functionalities

20 in peptides have been permethylated to generate a chemically transformed combinatorial library, is described by Ostresh et al. (1994, Proc. Natl. Acad. Sci. USA 91:11138-11142).

Screening the libraries can be accomplished by any of a variety of commonly known methods. See, e.g., the 25 following references, which disclose screening of peptide libraries: Parmley and Smith, 1989, Adv. Exp. Med. Biol.

251:215-218; Scott and Smith, 1990, Science 249:386-390; Fowlkes et al., 1992; BioTechniques 13:422-427; Oldenburg et al., 1992, Proc. Natl. Acad. Sci. USA 89:5393-5397; Yu et

- 30 al., 1994, Cell 76:933-945; Staudt et al., 1988, Science 241:577-580; Bock et al., 1992, Nature 355:564-566; Tuerk et al., 1992, Proc. Natl. Acad. Sci. USA 89:6988-6992; Ellington et al., 1992, Nature 355:850-852; U.S. Patent No. 5,096,815, U.S. Patent No. 5,223,409, and U.S. Patent No. 5,198,346, all
- 35 to Ladner et al.; Rebar and Pabo, 1993, Science 263:671-673; and PCT Publication No. WO 94/18318.



In a specific embodiment, screening can be carried out by contacting the library members with a lats protein (or nucleic acid or derivative) immobilized on a solid phase and harvesting those library members that bind to the protein (or nucleic acid or derivative). Examples of such screening methods, termed "panning" techniques are described by way of example in Parmley and Smith, 1988, Gene 73:305-318; Fowlkes et al., 1992, BioTechniques 13:422-427; PCT Publication No. WO 94/18318; and in references cited hereinabove.

In another embodiment, the two-hybrid system for selecting interacting proteins in yeast (Fields and Song, 1989, Nature 340:245-246; Chien et al., 1991, Proc. Natl. Acad. Sci. USA 88:9578-9582) can be used to identify molecules that specifically bind to a lats protein or derivative.

In addition, Drosophila can be used as a model system in order to detect genes that phenotypically interact with lats. For example, overexpression of lats in Drosophila eye leads to a smaller and rougher eye. Mutagenesis of the fly genome can be performed, followed by selecting flies in which the mutagenesis has resulted in suppression or enhancement of the small rough eye phenotype; the mutated genes in such flies are likely to encode proteins that interact/bind with lats.

25

## 5.15. <u>ANIMAL MODELS</u> The invention also provides animal models.

In one embodiment, animal models for diseases and disorders involving cell overproliferation (e.g., as described in Section 5.8.1) are provided. Such an animal can be initially produced by promoting homologous recombination between a lats gene in its chromosome and an exogenous lats gene that has been rendered biologically inactive (preferably by insertion of a heterologous sequence, e.g., an antibiotic resistance gene). In a preferred aspect, this homologous recombination is carried out by transforming embryo-derived stem (ES) cells with a vector containing the insertionally

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inactivated lats gene, such that homologous recombination occurs, followed by injecting the ES cells into a blastocyst, and implanting the blastocyst into a foster mother, followed by the birth of the chimeric animal ("knockout animal") in which a lats gene has been inactivated (see Capecchi, 1989, Science 244:1288-1292). The chimeric animal can be bred to produce additional knockout animals. Such animals can be mice, hamsters, sheep, pigs, cattle, etc., and are preferably non-human mammals. In a specific embodiment, a knockout nouse is produced.

Such knockout animals are expected to develop or be predisposed to developing diseases or disorders involving cell overproliferation (e.g., malignancy) and thus can have use as animal models of such diseases and disorders, e.g., to screen for or test molecules (e.g., potential anti-cancer therapeutics) for the ability to inhibit overproliferation (e.g., tumor formation) and thus treat or prevent such diseases or disorders.

In a different embodiment of the invention,

20 transgenic animals that have incorporated and express a
functional lats gene have use as animal models of diseases
and disorders involving deficiencies in cell proliferation or
in which cell proliferation is desired. Such animals can be
used to screen for or test molecules for the ability to

25 promote proliferation and thus treat or prevent such diseases
and disorders.

# 5.16. METHODS OF IDENTIFYING TUMOR SUPPRESSOR GENES AND OTHER GENES WITH IDENTIFIABLE PHENOTYPES

The invention also provides methods of identifying a tumor suppressor gene (or potential tumor suppressor gene) comprising identifying an overproliferation phenotype in a genetic mosaic, and isolating a gene that is mutated in cells exhibiting the overproliferation phenotype. The genetic mosaic is achieved by induction of somatic cells in an animal that is heterozygous for an induced mutation to become

homozygous for the mutation, at any desired developmental stage. The mutation can be induced by any known method, e.g., X-ray exposure or chemical mutation exposure or insertion of a transposable element (e.g., P-element). A genetic mosaic is produced by induction of homozygosity by mitotic recombination between homologous arms of both parental chromosomes, which is achieved using a site-specific recombination system [a sequence capable of expressing a site-specific recombinase; and its target sites (sequences at which the recombinase promotes recombination)], that have been inserted in the homozygous arms of both parental chromosomes. The target sites are preferably inserted close to the centromere on each chromosome arm (the closer to the centromere, the more preferred), so that mitotic

- 15 recombination events will result in cells being homozygous for the mutation located on the chromosome arm distal to the insertion of the target site. For example, an FLP recombinase can be used with FRT target sites; Cre recombinase can be used with lox target sites. The
- 20 recombinase coding sequence, used to express recombinase, preferably, but need not be, intrachromosomally situated. For at least one chromosome, the target sites are intrachromosomally inserted on the homologous arms of both parental (maternal and paternal) chromosomes.
- 25 The genetic mosaic can be an animal, e.g., mouse, hamster, sheep, pig, cow, *Drosophila*, etc., and is preferably a non-human mammal.

In a specific embodiment relating to the production of a non-human mammal that is a genetic mosaic, a recombinase 30 target site is introduced onto one arm of a chromosome in an embryo-derived stem cell (ES). The target site can be introduced into the cell by homologous recombination (by use of flanking sequences from the desired site of intrachromosomal integration) or by random integration 35 resulting from cell transformation (e.g., by transfection, electroporation), etc. This ES is then injected into a blastocyst, the blastocyst is implanted into a foster mother,

followed by birth of the recombinant animal. This mammal is bred to a wild-type female, to produce siblings. Siblings carrying the target site insertion are mated, and offspring carrying the target site on the homologous arms of both

- 5 parental chromosomes are isolated ("the target strain"). A target strain member is then mutagenized and mated with a non-mutagenized target strain member of the opposite sex (preferably also carrying a recombinant nucleic acid encoding and capable of expressing a recombinase that promotes
- 10 recombination at the target sites), to obtain a target strain member that is heterozygous for the mutation. Provision of the recombinase (by expression) in mitotically active cells of a developing animal or an adult animal promotes mitotic recombination between the homologous arms of the parental
- 15 chromosomes, resulting in a cell that is homozygous for the mutation. Cells that display a mutant phenotype by virtue of their being homozygous for the mutation are then detected, and the mutant gene can be genetically mapped by any known method, and can be isolated.
- In a *Drosophila* animal, a site-specific recombination system can be introduced by use of P-element-mediated insertions.

In one embodiment, target sites are introduced onto homologous arms of both of a set of parental chromosomes, for 25 one chromosome. In another embodiment, target sites are introduced onto homologous arms of both of a set of parental chromosomes, for a plurality of chromosomes.

The recombinase can be under the control of a constitutive (e.g., phosphorylated kinase promoter) or

30 inducible (e.g., heat shock promoter) or tissue-specific promoter. The recombinase can be expressed episomally (e.g., from a plasmid) or chromosomally. Once the recombination system is introduced into the animal, genetic mosaicism is produced by the activity of the recombinase (which promotes recombination at the target sites).

In a specific embodiment, an animal is used that contains a recombinant nucleic acid encoding an FLP

recombinase (Broach and Hicks, 1980, Cell 21:501-508) such that it is expressible by a cell of the animal, and intrachromosomal insertions of an FRT site on the homologous arms of both parental chromosomes; and genetic mosaicism is produced by inducing mitotic recombination between the FRT sites on the homologous chromosome arms after FLP recombinase expression (e.g., by heat shock, when expression of the FLP recombinase is under the control of a heat shock promoter).

In another specific embodiment, an animal is used

10 that contains a recombinant nucleic acid encoding a Cre
recombinase (Sauer and Henderson, 1988, Proc. Natl. Acad.
Sci. USA 85:5166-5170) such that it is expressible by a cell
of the animal, and intrachromosomal insertions of a lox site
on homologous arms of both parental chromosomes; and genetic

15 mosaicism is produced by inducing mitotic recombination
between the lox sites on the homologous chromosome arms after
Cre recombinase expression.

The animal may optionally further comprise intrachromosomal insertions of marker genes (comprising a sequence encoding a protein containing a reporter group such as an epitope tag), to facilitate confirmation and/or monitoring of recombination events. For example, in a non-human mammal, a marker gene (e.g., lacz) operably linked to a constitutive promoter can be inserted, on the same chromosome arm as that carrying the target site and the induced mutation.

In a specific embodiment, the overproliferation phenotype is the formation of overproliferated outgrowth tissue in a non-position-dependent fashion. In another specific embodiment, the overproliferation phenotype is the formation of a normal structure of larger than normal size.

The above-described genetic mosaics have uses not only in identifying tumor suppressor genes, but, more generally, in identifying genes with an identifiable

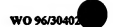
35 phenotype, i.e., those genes which in mutated form cause an observable mutant phenotype to be displayed in the genetic mosaic.

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In another embodiment, the invention provides a method of identifying genes with an observable mutant phenotype by use of human (or other animal) tissue culture cells that have incorporated a site-specific recombination 5 system such as described above. The site-specific recombination system can be introduced by methods such as described above, so as to introduce a recombinant source of recombinase and effect intrachromosomal insertions of the recombinase target sites on the homologous arms of both of a 10 set of parental chromosomes, for one or more chromosomes. a preferred aspect relating to this use of culture cells, the recombinase target sites are ligated to a selectable marker (e.g., an antibiotic resistance gene), and cells are obtained that have the target sites on each of the homologous 15 chromosome arms, by selecting under selection conditions of relatively high stringency (e.g., by increasing the antibiotic concentration in the cell medium), As with the use of genetic mosaics as described above, once mitotic recombination is induced between the target sites on the 20 homologous chromosome arms, one then identifies cells displaying a mutant phenotype, and recovers a gene mutated in cells exhibiting the mutant phenotype. For example, a potential tumor suppressor gene can be identified by isolating a gene that is mutated in cultured cells exhibiting 25 a transformed phenotype.

6. IDENTIFYING TUMOR SUPPRESSORS IN GENETIC MOSAICS: THE DROSOPHILA LATS GENE ENCODES A PUTATIVE PROTEIN KINASE

we have identified recessive overproliferation
mutations by screening and examining clones of mutant cells
in genetic mosaics of the fruitfly Drosophila melanogaster
(Fig. 1A). Flies that carry small groups of somatic cells
mutated for negative regulators of cell proliferation or
tumor suppressors are viable, yet the overproliferated mutant
tissues can be readily identifiable.



One way to generate mosaic animals is to induce mitotic recombination in developing heterozygous individuals (Fig. 1B). Recently, it was found that the site-specific recombination system from yeast, the FLP recombinase and its target site FRT, can be used to induce high frequency of mitotic recombination in Drosophila (Golic and Lindquist, 1989, Cell 59:499-509; Golic, 1991, Science 252:958-961). To produce and analyze genetic mosaics, a series of special Drosophila strains were constructed, containing the FLP/FRT recombination system on genetically marked chromosomes (Xu and Rubin, 1993, Development 117:1223-1237). Using these strains, high frequencies of mosaicism can be produced for more than 95% of the Drosophila genes. We have used these strains to identify overproliferation mutations in mosaic

Our results show that screening for overproliferation mutations in mosaic animals is a powerful way to identify negative regulators of cell proliferation and potential tumor suppressor genes. One of the identified genes, large tumor suppressor (lats), has been cloned, and encodes a predicted novel protein kinase. Mutations in lats cause dramatic overproliferation phenotypes and various developmental defects in both mosaic animals and homozygous mutants.

25

#### 6.1. MATERIALS AND METHODS

#### Genetics

rly stocks and crosses were grown on standard medium at 25°C unless otherwise indicated. The F1 mosaic screens were modified from the one described in Xu and Rubin (1993, Development 117:1223-1237) and in Xu and Harrison (1994, Methods in Cell Biology 44:655-682). Briefly, the F1 mosaic individuals were produced from three crosses:

Mutagenized y w hsFLP1; P[ry+; hs-neo; FRT]40A males were mated to the y w hsFLP1; P[ry+; y+]25F, P[mini-w+; hs-NM]31E, P[ry+; hs-neo; FRT]40A females. Mutagenized y w hsFLP1; P[ry+; hs-neo; FRT]42D males were mated to the y w hsFLP1;

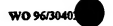


P[ry+; hs-neo; FRT]42D, P[ry+; y+]44B, P[mini-w+; hs-NM]46F/CyO females. Finally, mutagenized y w hsFLP1; P[ry+; hs-neo; FRT]82B males were mated to the y w hsFLP1; P[ry+; hs-neo; FRT]82B, P[mini-w+; hs-πM]87E, Sb<sup>635</sup>, P[ry+; y+]96E

- 5 females. The male parents were irradiated with X-rays (4000 r) and were removed from the crosses after four days of mating. The eggs from the crosses were collected for every 12 hours and aged for another 30 hours before being incubated in a 38°C water bath for 60 minutes. The F<sub>i</sub> animals were then
- 10 returned to normal culture conditions until eclosion. About 25,000  $F_1$  adults from these crosses were examined. Each P-induced lethal mutation was recombined onto one of the FRT-carrying arms using the  $neo^R$  and w double selection as described in Xu and Harrison (1994, Methods in Cell Biology
- 15 44:655-682) before examining its clonal phenotype.

The lats' mutation was meiotically mapped to the right of claret. It was further localized to the 100A1-5 region since it complemented Df(3R)tll'(100A2-5; 100C2-3) and failed to complement  $Df(3R)tll'^{\mu\nu}(100A1-2; 100B4-5)$  and

- 20 Df (3R) t11<sup>20</sup>(100A1-3; 100B1-2). A saturation genetic screen had previously been performed for this interval, and three lethal complementation groups, 1(3)100Aa, 1(3)100Ab and the zfh-1, were isolated (Lai et al., 1993, Proc. Natl. Acad. Sci. USA 90:4122-4126). The lats<sup>11</sup> mutation failed to
- 25 complement the EMS-induced mutations in 1(3)100Aa (lats<sup>al-al5</sup>), but complement mutations in 1(3)100Ab and zfh-1. The clonal phenotypes were examined for lats<sup>xl,Pl,al,a2,ab and al0</sup> induced either with the FLP/FRT-marker system or X-ray irradiation.
- The lats<sup>PI</sup> allele was recovered from a mosaic male
  produced from the cross of y w hsFLP1; P[ry\*; hs-neo; FRT]82B
  x y w P[lacZ; w\*]5; P[ry\*; hs-neo; FRT]82B/delta2-3, Sb. The
  mutant chromosome was cleaned up before performing
  complementation tests and an excision screen (Robertson et
  al., 1988, Genetics 118:461-470). Two hundred and fifteen
  excision lines were established that had lost the w\* gene in
  the P[lacZ; w\*] element (Bier et al., 1989, Genes Dev.



3:1273-1287). In about 50% of these lines, the pupal
lethality had been reverted completely to wild type,
indicating the mutant phenotype is caused by the P-element
insertion. Pive lines were found to cause lethality at late
sembryonic and/or early first instar larval stages. The
remaining lines were found to cause lethality at larval and
pupal stages or to produce viable mutant animals. All of
these mutant excision lines (except one which is located
outside the 100A1-5 region) failed to complement lats"and
lats", but do complement mutations in the zfh-1 and 1(3)100Ab
loci.

The insert in lats cDNA A2 was cloned into the pCaSpeR-hs vector (Thummel and Pirrotta, 1992, Drosophila Inform. Service 71:150) for germ line transformation. Three of the transformed lines were tested and were able to rescue the lethality of the lats lats lats lats lats lats and lats and lats after one hour heat shock for every 24 hours during larval and pupal development.

#### 20 Histology

Fixation and sectioning (2 mm) of adult Drosophila tissues were performed as described (Tomlinson and Ready, 1987, Dev. Biol. 123:264-275). Scanning electron microscopy was performed as described (Xu and Artavanis-Tsakonas, 1990, 25 Genetics 126:665-677).

#### Nucleic Acid Manipulation

region was obtained from the Berkeley Drosophila Genome

30 Center (personal communication; Hartl et al., 1994, Proc.
Natl. Acad. Sci. USA 91:6824-6829). DNA fragments from this
P1 clone and genomic DNA obtained by plasmid rescue from the
lats<sup>P1</sup> mutant (Bier et al., 1989, Genes Dev. 3:1273-1287) were
used to isolate several overlapping cosmids including CLT-52

35 from the genomic library prepared by J. Tamkun. Genomic DNA
from +7.5 (BglII) to -4.2 (EcoRI; Fig. 3) was used to screen
a total imaginal disc cDNA library prepared by A. Cowman.



Screening approximately 2 million phage yielded three groups of cDNAs (five lats cDNAs; fifteen T1 cDNAs; fourteen T2 cDNAs). The sizes of the inserts in the lats cDNAs are as follows: 5.6 kb in A2; 5.1 kb in B1; 1.1 kb in 9 and 4; and 5 0.9 kb in B3.

Genomic DNA from lats<sup>1</sup>/TM6B, lats<sup>2-15</sup>/TM6B, lats<sup>2-15</sup>/TM6B, lats<sup>2-17</sup>/TM6B, lats<sup>2-17</sup>/TM6B, lats<sup>2-18</sup>/TM6B, lats<sup>2-18</sup>/TM6B flies was digested with a combination of the EcoRI, BamHI, BglII and XhoI restriction enzymes for Southern analysis.

#### DNA Sequencing

DNA sequence was determined by the dideoxy chain termination method (Sanger et al., 1977, Proc. Natl. Acad. 15 Sci. USA 74:5463-5467) using Tag polymerase (Perkin Elmer) and Sequenase (U.S. Biochemical Corp.). The sequences of lats cDNAs were determined from both strands using templates generated from plasmids containing EcoRI fragments inserted into the pBlueScriptII vector. Templates generated from 20 DNase 1 deletion subclones were also used. The complete sequences of cDNAs A2 and 9 were determined; partial sequences were determined for cDNAs B1 and 4. Templates of genomic DNA were generated from plasmids containing EcoRI fragments and were sequenced on one strand using synthetic 25 oligonucleotide primers. Mutant DNA from the lats allele was amplified with PCR reactions using synthetic oligonucleotide primers and cloned in the pBlueScript II vector for sequencing.

#### 30

#### 6.2. RESULTS

Screening for Overproliferation Mutations in Mosaic Animals

We have screened individuals carrying clones of cells that were homozygous for either X-ray or P-element induced mutations for overproliferation phenotypes. (Fig. 1B; Materials and Methods). Two types of overproliferation phenotypes were sought: a) Clones of mutant cells formed

overproliferated, outgrowth tissues in a non-positiondependent fashion; b) Clones of mutant cells formed normal structures, but proliferated faster than wild-type cells such that the sizes of the mutant clones were larger than their wt 5 twin-spot clones. Three independent mutations were identified that caused the first type of phenotype (Fig. 2A-2E). A mutation which was allelic to one of the original mutations was later found to cause the second type of phenotype (see below). All three mutations in the first 10 class caused embryonic and/or early larval lethality and they represented single alleles of different loci since they had different chromosome locations. One of them was identified among 215 randomly chosen lethal mutations in which each were caused by a P-element insertion in a different essential gene 15 (Karpen and Spradling, 1992, Genetics 132:737-753; Berkeley Drosophila Genome Center, personal communication). addition to these overproliferation mutations, one P-induced mutation was found to cause both unpatterned outgrowth and duplications of patterned structures in mosaic animals, 20 suggesting that this mutation may not directly affect cell proliferation.

The lats Locus Is Defined by a Single Complementation Group of Mutations
That Cause Defects Throughout Development

overproliferation. One mutation (lats") produced much more dramatic overproliferated clones than the ones produced by the other mutations (Fig. 2A, 2B). The lats mutant clones induced in first instar larvae can be as large as 1/5 of the body size. Tumorous outgrowth caused by lats" was found in all the tissues that had been examined including eyes, legs, wings, heads, notums, antenna, and abdominal cuticles. The lats" mutation was genetically mapped in the 100A1-5 region and the locus was further defined by a single complementation group of over fifty alleles including mutations induced by



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X-ray, EMS, P-element insertion and imprecise excision of the P-element (Table 2; Materials and Methods).

#### TABLE 2

5

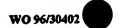
#### The alleles of the lats locus

	Alleles	Phenotypes of homozygous animals	Phenotypes of mutant clones	Representative alleles	No. of alleles
10	Strong	Late embryonic and early 1st instar larval lethal	Large outgrowth	lats", lats", lats"	14
	Medium	Late larval and pupal lethal, normal size of animals	Large outgrowth	lats <sup>e</sup> , lats <sup>d24</sup>	16
15	Weak	Pupal lethal, giant animals	Large outgrowth	lats <sup>ex-1</sup>	3
		Semi-viable and viable: rough eye outgrowth on head, wing held-out, sterile	Mutant clones larger or normal in size	lats <sup>alv</sup> , lats <sup>,53.</sup>	

The various alleles of the lats gene are classified into three main groups as indicated in the left column. Their phenotypes, displayed in either homozygous mutant animals or clones of mutant cells in mosaic animals, are listed in the next two columns respectively. For a given viable or semi-viable allele, the homozygous mutant animals display one, two, three, or all four of the listed phenotypes. Representative alleles and the numbers of alleles for each group are given in the two right columns. The origins of these alleles are described in the Material and Methods.

chromosome into wild type, indicating the P-element insertion is responsible for the mutant phenotype. Furthermore, five of the imprecise excision lines caused late embryonic and early larval lethality which were stronger than the pupal lethality phenotype caused by the lats<sup>PI</sup> mutation. These five excision lines failed to complement lats<sup>II</sup>, but complemented the mutations in two other complementation groups (1(3)100Ab and zfh-1) in the 100A1-5 region, indicating that these two genes were not affected by the excision alleles.

The lats alleles can be classified into three main groups (Table 2). Strong alleles caused homozygous animals to die at a late embryonic stage or shortly after hatching



with no obvious cuticular defect. Mutations in the group of medium alleles cause lethality at different times in larval and pupal development. This group was further divided into two subgroups because three of the excision alleles not only caused pupal lethality, but the sizes of the homozygous mutant animals were also significantly larger than wt animals (Fig. 2C). The weak mutations caused either one or a combination of the following phenotypes: held out wings with broadened blades, rough eye with ventral outgrowth, outgrowth on the dorsal-anterior region of the head and partial to

complete sterility (Table 2). Proliferation defects were observed in both mutant clones in mosaic animals and homozygous mutants. Clones of cells on the head that were homozygous for strong or medium 15 alleles formed unpatterned, overproliferated tissues with many lobes or folds. The mutant cells seemed to be "budding out" of the surface to form new proliferation centers or lobes (Fig. 2A, 2F, 2H). The sizes and the shapes of these mutant cells were very irregular. Cells several times larger 20 than their neighbors were often seen in mutant clones, indicating problematic cell division (Fig. 2F, 2G). Furthermore, lats mutant clones behaved differently from clones mutant for the previously identified Drosophila tumor suppressor genes such as dlg, lgl and hyd. The dlg, lgl or 25 hyd mutant cells proliferated slower than wt cells and thus, the mutant clones induced in first instar larvae were competed away during growth and did not form detectable clones in the adults (Bryant, 1987, Experimental and genetic analysis of growth and cell proliferation in Drosophila

- 30 imaginal discs, in "Genetic Regulation of Development," A.R. Liss, New York, pp. 339-372; Woods and Bryant, 1989; Dev. Biol. 134:222-235; Mansfield et al., 1994, Dev. Biol. 165:507-526; Allen Shearn, personal communication). In contrast, the lats mutant clones induced at similar
- 35 developmental stages formed dramatic overproliferated tissues, suggesting the mutant cells proliferated faster than wt cells. Consistent with this notion, clones of cells

mutant for a weak lats allele (latsalo) produced normal looking tissues, but the mutant clones were significantly larger than their wt twin-spot clones. In homozygous animals, the imaginal discs and the central nervous system in many of the 5 pupal lethal mutants were dramatically overproliferated (Fig. 2D, 2E). The discs lost the single layer of epithelial structure and formed multi-layer, deformed tissues. overproliferation phenotype was not caused by prevention of differentiation. Cells in the overproliferated mutant clones 10 on the body differentiated and produced bristles and hairs, although the morphologies of these structures were not wild type (Fig. 2I-2L). Careful examination of multiple mutant clones confirmed that lats caused mutant cells (w cells in the eye, y bristles and enlarged-base hairs on the body) to 15 overproliferate and did not affect the surrounding wt tissues. Finally, the frequency of overproliferated clones was similar to wt clonal frequency induced with the same FRT element, indicating that loss of the lats function alone is sufficient to initiate the overproliferation process.

#### Cloning of the lats Gene

20

Genomic DNA from the 100A1-5 region was isolated using probes mapped to this region (Materials and Methods). restriction map of the relevant genomic region is illustrated 25 in Figure 3. Genomic DNA flanking the P-insertion site (+7.5 to -4.2) was used to screen a total imaginal disc cDNA library. A group of cDNAs corresponding to a 5.7 kb transcript (lats) was found to contain sequence from the region where the P-element was inserted (Fig. 3). Two other 30 groups of cDNAs were also isolated (T1 and T2). The 5.7 kb transcript was located in an intron of the T1 gene (Fig. 3). The intron-exon structure of the 5.7 transcription unit was determined by Southern and sequence analysis of the cDNA clones and the corresponding genomic DNA (Materials and 35 Methods). The zfh-1 gene was found to be located at the left side of the 5.7 kb transcription unit (Fig. 3; Fortini et al., 1991, Mechanisms of Development 34:113-122).

In addition to lats<sup>PI</sup>, genomic DNA from the five strong excision alleles was analyzed. All of them deleted exon sequences from the 5.7 kb transcript and, in addition, three of them also deleted sequences in the next transcript

- 5 (T2; Fig. 3). Furthermore, DNA from the X-ray and EMS induced mutants was analyzed with cDNA probes made from the 5.7 kb, T2 and T1 transcripts. In two cases alterations were detected in the 5.7 kb transcription unit: a 0.4 kb and a 0.3 kb deletions associated with lats<sup>al</sup> and lats<sup>al</sup>, respectively
- 10 (Fig. 3). The 446 bp deletion in lats<sup>al</sup> was revealed by sequencing. It removed codons 92 to 238 of the open reading frame and caused a frame shift from codon 239 (Fig. 5). Finally, transformants containing a cDNA corresponding to the 5.7 transcript driving by the hsp70 promoter rescued the
- 15 lethality of both strong and medium lats alleles. These findings indicate that the 5.7 kb transcription unit which correspond to the lats gene and strong lats alleles including lats. were either amorphic or nearly amorphic alleles.

## 20 The lats Gene Encodes a Putative Protein-Serine/Threonine Kinase

The 5.7 kb lats transcript was detected throughout development (Fig. 4) and in both adult males and females (data not shown). In addition, probes from the 5.7 kb transcript also detected a second transcript, which is about 1 kb shorter (4.7 kb), in young embryos (0-4 hrs; Fig. 4) and

- in adult males and females. Northern analysis showed there was more maternally deposited 4.7 kb transcripts than 5.7 kb transcripts in young embryos (0-2 hrs; Fig. 4). The 5.7 kb transcript became the dominant message at the embryonic stage (4-6 hrs), known to have zygotic gene expression (Fig. 4).
- No effort was made to isolate cDNA clones corresponding to the 4.7 kb transcript; thus the exact sequence of this short transcript is not known. However, a polyadenylation signal consensus sequence was found at nucleotide position 4655 4660 in the 5.7 kb transcript and in the corresponding
- 4660 in the 5.7 kb transcript and in the corresponding genomic DNA (Fig. 5) and a 0.51 kb probe from the 3' end of



the 5.7 kb transcript did not hybridize to the 4.7 kb transcript while a 1 kb probe from the 5' untranslated region of the 5.7 kb transcript hybridized to both the 5.7 kb and 4.7 kb transcripts. This suggests that the 4.7 kb transcript

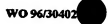
- 5 may be a truncated version of the 5.7 kb transcript. The genomic and cDNA sequence corresponding to the 5.7 kb transcript was determined (Materials and Methods). The entire 5720 bp cDNA sequence, which is interrupted by seven introns, and the putative lats product (lats), deduced from
- 10 the long open reading frame, are illustrated in Figure 5. An interesting feature of the 5.7 kb transcript is the existence of a 141 bp segment located in the 3' untranslated region (Fig. 5), which is identical to the first 141 bp of the 5' untranslated region of the class I transcript from the
- 15 Drosophila phospholipase C gene, plc-21 (Shortridge et al., 1991, J. Biol. Chem. 266:12474-12480). The functional significance of this sequence motif is unknown. It could be a regulatory target sequence that is shared by both genes.

There are 34 differences between the lats cDNA and genomic sequences and 31 of them do not affect the deduced amino acid sequence. In the remaining three differences, one changes the serine 206 in cDNA into a cysteine. The second change in the genomic sequence adds an additional glutamine in the poly-glutamine opa repeat (Fig. 6; Wharton et al.,

25 1985, Cell 40:55-62). The third is the addition of a fifteen bp sequence in the genomic DNA after the nucleotide 2644 of the cDNA. This sequence could be translated into another copy of the Arg-Glu-Arg-Asp-Gln (part of SEQ ID NO:2) peptide. However, this sequence is not present in the two independent cDNA clones that were sequenced.

The predicted lats product contains 1099 amino acid residues. The kinase domain of lats is more similar to protein-serine/threonine kinases than to protein-tyrosine kinases, especially in the sequences of the domains VI and

35 VIII defined by Hanks et al. (1988, Science 241:42-52); protein-serine/threonine kinase consensus in domain VI: Asp-Leu-Lys-Pro-Glu-Asn (SEQ ID NO:9). Lats sequence in domain



VI: Arg-Asp-Ile-Lys-Pro-Asp-Asn (836-842) (part of SEQ ID NO:2); protein-serine/threonine kinase consensus in domain VIII: Gly-Thr/Ser-X-X-Tyr/Phe-X-Ala-Pro-Glu (SEQ ID NO:10). Lats sequence in domain VIII: Gly-Thr-Pro-

- 5 Asn-Tyr-Ile-Ala-Pro-Glu (917-925) (part of SEQ ID NO:2). The C-terminal half of lats shares extensive sequence similarity with a group of six proteins including the Dbf20 and Dbf2 cell cycle protein-ser/thr kinases from Saccharomyces cerevisiae (Johnston et al., 1990, Mol. Cell. Biol. 10:1358-
- 10 1366; Toyn et al., 1991, Gene 104:63-70; Toyn and Johnston, 1994, EMBO J. 13:1103-1113), and the COT-1 putative protein kinase from Neurospora crassa (Yarden et al., 1992; EMBO J. 11:2159-2166) (Fig. 6A, 6B). The sequence similarity between the kinase domains of lats and these proteins (39-49%
- observed between the different subgroups of protein-ser/thr kinases (20-25% identity; Hanks et al., 1988, Science 241:42-52). However, there is an insertion of about 40 amino acid residues within the kinase domains of these proteins,
- 20 sharing little sequence similarity (denoted by a black bar in Fig. 6B). The human myotonic dystrophy protein kinases (MDPK) also have significant similarity with the C-terminal region of lats (Brook et al., 1992, Cell 68:799-808; Fu et al., 1993; Science 260:235-238, Mahadevan et al., 1993, Hum.
- 25 Mol. Genet. 2:299-304), but their kinase domains do not contain this ~40 amino acid insertion. In addition, lats and these proteins also share significant levels of sequence similarity in the two regions (each contains ~100-150 amino acids) flanking the kinase domain (20-28% identity; Fig. 6A,
- 30 6B). In the case of Dbf20, its entire sequence except for the 20 C-terminal most residues can be aligned with lats, indicating lats is a close relative of Dbf20. A polyglutamine opa repeat is located near the middle of the protein (Fig. 5; Wharton et al., 1985, Cell 40:55-62). The
- 35 N-terminal half of lats contains many short homopolymeric runs including poly-proline which makes up about 15% of the residues. At least one of the proline-rich stretches closely



matches the consensus of SH3-binding sites (Fig. 3B; Ren et al., 1993, Science 259:1157-1161), raising the possibility that it may interact with SH3-containing proteins. No putative signal sequence appears in the lats protein, 5 indicating that it is an intracellular protein.

#### 6.3. DISCUSSION

Screening for Mutations in Mosaic Animals to Identify and Study Potential Tumor Suppressors

10 The comparison between mosaic flies and tumor patients is simplistic yet useful. Tumor patients contain wt tumor suppressor genes in most of their cells and only small groups of cells sustain mutations in tumor suppressors. searched for recessive overproliferation mutations in mosaic animals. Flies that carry somatic cells mutated for tumor suppressors or negative regulators of cell proliferation are viable, yet the overproliferation mutant phenotype is readily identifiable. Therefore, mosaic flies, which are in a fashion analogous to tumor patients, provide a mean to screen for potential tumor suppressors. Three overproliferation mutations were identified in our screen. They were not identified as "interesting" mutations in screens for embryonic lethal mutations. Identifying overproliferation mutations in homozygous mutant larvae and pupae is not only biased against embryonic lethals, but also laborious, since it requires establishment of individual lines before examining the potential phenotypes. Further screens for overproliferation mutations in mosaic animals will allow us to identify other important players in pathways that negatively regulate cell proliferation.

The overproliferation phenotypes that we observed were caused by loss of function in a single gene. In humans, it was suggested that most retinoblastomas are caused by defects in a single tumor suppressor (Knudson, 1971, Proc. Natl. Acad. Sci. USA 68:820-823). On the other hand, evidence indicates that tumorigenesis in other human tissues (e.g.,

colon cancer) is a multistep process which involves inactivation of more than one gene (Fearon and Vogelstein, 1990, Cell 61:759-767; Vogelstein and Kinzler, 1993, Trends Genet. 9:138-141). Overproliferation caused by defects in multiple genes is unlikely to be detected in our screens unless these genes are located on the same chromosome arm. To identify this type of gene, one could perform a modified mosaic screen which induces clones of cells to become homozygous for more than one mutagenized chromosome arm.

10

## lats Affects Many Tissues Throughout Development

The lats gene is genetically defined by a single complementation group that consists of various alleles causing a wide range of defects. Different alleles were

- 15 found to cause lethality at almost every stage during development: embryo, early larvae, late larvae, early pupae, late pupae and pharate-adult. The embryonic lethality occurs in the pharate first instar stage. The early embryonic requirements for lats could well be masked by the wt products
- 20 that are maternally deposited in the egg. Weak lats alleles produce viable animals with phenotypes ranging from rough eye to sterility. The lats transcripts were detected throughout development up to adult stage, consistent with the observation that lats mutants affect all these stages.
- 25 Although mutations at lats cause many defects, affecting cell proliferation could cause most of the phenotypes including overproliferation in mutant clones, lethality at the various stages, tissue overproliferation on the head, broadened wing blade, and sterility in homozygous mutants. However,
- 30 phenotypes such as extra cuticle deposits and malformed bristles and hairs are evidence of defects in differentiation.

The different behavior of the lats mutant clones and clones mutant for other previously identified Drosophila

35 tumor suppressors is interesting. Cells mutant for dlg, lgl or hyd seem to fail to receive growth regulation signals. They proliferated slower than wt cells during larval stages



when the cells were instructed to proliferate, and they failed to terminate proliferation in late larval and pupal stages when the wt cells have ceased proliferation. On the other hand, the lats mutant clones induced during the larval stages were overproliferated, and later the mutant cells on the body were differentiated to form adult cuticular structures. Thus, lats could be a negative regulator that monitors the rate of proliferation.

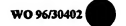
The lats gene is located in a complex region. The 5'

10 end of the lats 5.7 kb transcript (cDNA) is only about 550 bp
away from the T2 transcript and its 3' end is about 1.5 kb
away from the zfh-1 transcript. Furthermore, all three of
these closely located transcripts are located in an intron of
the T1 transcription unit. Thus, a sizable deletion in the
15 5.7 kb transcription unit could affect the function of any of
the genes in the region, which makes it difficult to
determine which transcript is responsible for the lats
phenotype. The fact that P-element transform lines carrying
a cDNA from the 5.7 kb transcript under the hsp70 promoter
20 rescued all types of lats alleles demonstrated that the 5.7
kb transcription unit is the lats gene.

The lats Putative Protein-Ser/Thr Kinase Shares Homology With Proteins That Are Involved in Regulation of Cell Cycle and Growth in Budding Yeast and Neurospora

All 11 subdomains of the kinase domain that are found in previously identified protein kinases (Hanks et al., 1988, Science 241:42-52) are conserved in lats. This predicts that lats is a protein kinase. Furthermore, the sequence

- 30 comparisons suggest lats to be a ser/thr kinase as the lats kinase domain is more similar to protein-ser/thr kinases than to protein-tyr kinases. The C-terminal half of lats shares extensive sequence similarity with a group of six proteins. Mutations are known for three of these genes and in each case
- 35 they affect either cell cycle or growth. The cot-1 (colonial temperature sensitive-1) gene of Neurospora was identified by a temperature sensitive mutant that causes compact colony



growth (Mitchell and Mitchell, 1954, Proc. Natl. Acad. Sci. USA 40:436-440; Galsworthy, 1966, Diss. Abstr. 26:6348). Wild-type filamentous ascomycete Neurospora grows on solid media by continuous hyphal elongation and branching to form

- 5 spreading colonies. Strains lacking functional cot-1 gene are viable, but their hyphae branch extensively, resulting in compact colonial growth (Yarden et al., 1992, EMBO J. 11:2159-2166). This extensive branching phenotype is somewhat similar to the growth property of the lats mutant
- 10 clones: the lats mutant cells continue to "bud" out of the surface to form new proliferation lobes. Another homologous gene, the DBF2 gene of the budding yeast, was identified in a genetic screen for mutations causing defects in DNA synthesis (Johnston and Thomas, 1982, Mol. Gen. Genet. 186:439-444).
- 15 The temperature sensitive alleles of DBF2 were found to both delay the initiation of S phase and also to arrest the cell cycle during nuclear division (Johnston et al., 1990, Mol. Cell. Biol. 10:1358-1366). The DBF20 gene was identified through cross hybridization with DBF2 DNA (Toyn et al., 1991,
- 20 Gene 104:63-70). Strains carrying deletions for either DBF2 or DBF20 are viable; however, deleting both genes in the same strain causes lethality. The kinase activities of both proteins have been shown to be specific for serine/threonine residues and are regulated during the cell cycle (Toyn and
- 25 Johnston, 1994, EMBO J. 13:1103-1113). In the case of Dbf20, its entire sequence except the 20 most C-terminal residues can be aligned with lats. The mutant phenotype of lats and its sequence homology with the cell cycle protein kinases is consistent with the notion that lats might be directly
- and involved in regulation of the cell cycle. The N-terminal half of lats contains many proline-rich stretches and at least one of them closely matches the consensus sequence of SH3 binding sites (Ren et al., 1993, Science 259:1157-1161), raising the possibility that this region could be a
- 35 regulatory domain for the lats kinase, which binds to SH3 domain-containing proteins.



In recent years, many protein kinases have been identified to be involved in regulation of the cell cycle and cell proliferation. While Weel is an inhibitor of the Cdc2 kinase (Russell and Nurse, 1987, Cell 49:559-567;

- 5 Featherstone and Russell, 1991, Nature 349:808-811), all other previously identified protein kinases are positive regulators of cell proliferation. They are either required for completion of the cell cycle or for signalling cells to proliferate. Lats is the first predicted protein-ser/thr
- 10 kinase that has been shown to cause overproliferation when its function is removed. Studies of lats and other overproliferation mutations in Drosophila will provide a better understanding of how cell proliferation is regulated during development and how mutations could lead to abnormal 15 growth.

## 7. ISOLATION AND CHARACTERIZATION OF MAMMALIAN LATS HOMOLOGS

As described herein, we have cloned and sequenced both mouse and human lats homologs.

7.1. <u>ISOLATION AND CHARACTERIZATION OF MOUSE LATS HOMOLOGS</u>
CDNA clones for two different lats homologs in mice were obtained as follows.

25 Screening of Mouse Homologs:

Probe:

A 2.2 kb BamHI fragment containing the kinase domain of the *Drosophila lats* gene was labeled with <sup>32</sup>P by random labeling

Library:

Newborn mouse brain lambda ZAP cDNA library from Stratagene

Hybridization

Condition:

45°C, overnight in 6x SSC

5x Denhart's

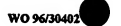
0.5% SDS (sodium dodecyl

sulfate)

100 μg/ml salmon sperm DNA

35 Wash: 50°C, 30 min. x 4, in 2x

2x SSC SDS



Results:

Three positive clones were identified. (M41 clone for the m-lats gene, and M51 and M31 clones for the m-lats2 gene.)

Two different mouse lats homologs, termed m-lats and m-lats2, respectively, were isolated and sequenced. the m-lats and m-lats2 clones are missing a small amount of the 5' end of their respective genes. The cDNA sequence (SEQ ID No:5) and deduced protein sequence (SEQ ID No:6) of m-lats are shown in Figure 7. The cDNA sequence (SEQ ID NO:7) and deduced protein sequence (SEQ ID NO:8) of m-lats2 are shown in Figure 8.

Portions of both the m-lats and m-lats2 cDNAs were used as probes to screen a mouse genomic library, under standard hybridization conditions. Genomic clones for both m-lats and m-lats2 have been isolated that contain most of the coding regions of these genes.

#### ISOLATION AND CHARACTERIZATION 7.2. OF HUMAN LATS HOMOLOGS

cDNA clones for at least one human lats homolog were obtained as follows.

Screening of Human Homologs (moderately stringent conditions):

Probe:

A 2.1 kb PstI fragment containing the kinase domain of the m-lats gene was labeled with 32P by random labeling

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Library:

Fetal human brain lambda gt10 cDNA library from Clontech

Hybridization

Condition:

55°C, overnight in 6x SSC

> Denhart's 5x

SDS 30 0.5%

salmon sperm DNA  $100 \mu g/ml$ 

Wash:

SSC 60°C, 30 min. x 2, in 1x

0.1% SDS

Results:

About 20 positive clones were identified for the h-lats gene.

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One human lats homolog, termed h-lats, was isolated and sequenced. The cDNA sequence (SEQ ID NO:3) and deduced

protein sequence (SEQ ID NO:4) of h-lats are shown in Figure
9. The deduced protein sequence is full-length. The complete coding sequence of the h-lats cDNA was inserted into a bacterial cloning vector (derived from Bluescript (KS)5 vector; Stratagene) to form plasmid pBS(KS)-h-lats (Fig. 10). The total size of pBS(KS)-h-lats is 6.96 kb.

A h-lats cDNA fragment was used as a probe under conditions of moderate stringency to screen a human genomic cosmid library. Genomic h-lats clones were isolated. Over 10 70 kb of the genomic h-lats sequence has been isolated; the isolated sequences include all of the h-lats coding sequence (but not all the exon sequences).

An m-lats2 cDNA fragment was used as a probe to screen a human genomic phage library under the conditions

15 described above, except that hybridization was carried out at 50°C, and washing was carried out at 55°C with 2X SSC, 0.1% SDS. Two genomic h-lats clones have been isolated that specifically hybridize to m-lats2 cDNA probes and do not hybridize to m-lats and h-lats cDNA probes.

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## 8. CONSERVATION OF SEQUENCES AND DOMAIN STRUCTURE AMONG LATS HOMOLOGS OF DIFFERENT SPECIES

Comparison of the sequences of Drosophila lats, h-lats, m-lats, and m-lats2 showed a startlingly high degree of sequence conservation, both overall and within domains of the lats protein. An alignment of the h-lats (SEQ ID NO:4) and m-lats (SEQ ID NO:6) protein sequences is shown in Figure 11. The overall amino acid sequence identity between h-lats and m-lats is 93%. An alignment of the h-lats (SEQ ID NO:4) and m-lats2 (SEQ ID NO:8) protein sequences is shown in Figure 12.

Homologous domains (i.e., domains conserved)
between the different lats homologs were identified. Figure
13 presents an alignment of the h-lats protein sequence (SEQ
ID NO: 4) and the Drosophila lats protein sequence (SEQ ID
NO:2), and indicates the domains identified as conserved
among the lats homologs from the various species.

The identified domains were as follows:

(1) Lats C-terminal domain 3 (LCD3)

The last three amino acids (VYV) are completely conserved in all four homologs including *Drosophila* lats, h-lats, m-lats, and m-lats2.

(2) Lats C-terminal domain 2 (LCD2)

h-lats 1077-1086
Drosophila lats 1075-1084

- This domain is completely conserved in all four homologs including Drosophila lats, h-lats, m-lats, and m-lats2 (10/10 identical residues).
  - (3) Lats C-terminal domain 1 (LCD1)

amino acid residues

h-lats 1032-1043 Drosophila lats 1035-1047

This domain is completely conserved among

Drosophila lats, h-lats, and m-lats (12/12 identical),

and is highly conserved between any of the foregoing and

m-lats2 (11/12 identical).

(4) Kinase domain

h-lats

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amino acid residues 703-1014 s 711-1018

This domain is highly conserved among the four homologs (76% identical between Drosophila lats and h-lats; 99% identical between h-lats and m-lats; 83% identical between h-lats and m-lats2).

A potential phosphorylation residue in *Drosophila* lats and the mammalian homologs that could lead to the activation of the lats kinases after phosphorylation was identified.

Activities of protein kinases are often regulated by varying the phosphorylation state of specific serine, threonine, and tyrosine residues. Phosphorylation of a serine or threonine within twenty residues upstream of



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an Ala-Pro-Glu consensus in subdomain eight of the kinase domain, is often required for catalytic activities of many protein-ser/thr kinases (Hanks et al., 1988, Science 241:42-52). For example, Thr167 and Thr197 are phosphorylated in Cdc2 of fission yeast and in the cardiac muscle adenosine 3',5'-phosphate dependent protein kinase, respectively (Ducommun et al., 1991, EMBO J. 10:3311-3319; Gould et al., 1991, EMBO J. 10:3297-3309; Shoji et al., 1983, Biochem.

22:3702-3709). A ser residue in a similar position of the lats kinase domain is conserved in *Drosophila* lats, h-lats, m-lats, and m-lats2 (Ser914 in *Drosophila* lats; Ser909 in h-lats). Thus, the activities of *Drosophila* lats and its mammalian homologs may be regulated by phosphorylation of this ser residue.

(5) Lats flanking domain (LFD)

amino acid residues h-lats 607-702 Drosophila lats 612-710

LFD is a domain that flanks and is amino-terminal to the kinase domain. This domain is highly conserved between *Drosophila* lats and h-lats (68% identical) and is also highly conserved between h-lats and m-lats2 (71% identical). This domain is completely conserved between h-lats and m-lats (100% identical).

(6) Lats split domain 1 (LSD1)

LSD1 Drosophila-lats 365-392 LSD1 anterior (LSD1a) h-lats 328-334 LSD1 posterior (LSD1p) h-lats 498-518

Certain lats domains have been termed split domains because the amino- (anterior) and carboxy- (posterior) portions of the domain appear separated from each other in at least one of the lats homologs. Split domains may constitute discontinuous binding/functional regions (e.g., brought together by tertiary structure). The LSD1a subdomain is completely conserved among Drosophila

lats, h-lats, and m-lats (7/7 identical), and is not conserved in m-lats. The LSD1p subdomain is conserved between the four homologs (14/21 identical among Drosophila lats, h-lats, and m-lats; 13/21 identical between h-lats and m-lats2). The LSD1a and LSD1p subdomains are adjacent to each other in Drosophila lats and are separated in the mammalian homologs.

(7) Lats split domain 2 (LSD2)

amino acid residues

LSD2 Drosophila lats 536-544

LSD2 anterior (LSD2a) h-lats 28-31

LSD2 posterior (LSD2p) h-lats 555-559

Both the LSD2a and LSD2p subdomains are completely conserved among the four homologs. However, the two subdomains are adjacent to each other in *Drosophila* lats and are separated in the mammalian homologs.

(8) Putative SH3-binding domain (SH3-binding)

amino acid residues h-lats 247-268 Drosophila lats 196-217

This domain is highly conserved among *Drosophila* lats, h-lats, and m-lats (10/22 identical), and does not exist in m-lats2.

The opa domain does not appear in the mammalian lats homologs.

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## 9. FUNCTIONAL INTERCHANGEABILITY OF THE HUMAN AND DROSOPHILA LATS HOMOLOGS

9.1. OVEREXPRESSION OF HUMAN LATS OR OF DROSOPHILA LATS CAUSES A SMALLER, ROUGH EYE IN DROSOPHILA

Overexpression of lats and h-lats in the developing Drosophila eye was carried out. The Drosophila lats cDNA and h-lats cDNA were each cloned into the pGMR P-element vector. This vector was constructed by Bruce Hay and Gerald M. Rubin at the University of California at Berkeley, and will direct the expression of a cDNA of interest in the posterior region of the developing third instar larval eye imaging disc of

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Drosophila. Ten independent transformant lines for each of the pGMR-lats and pGMR-h-lats constructs were generated. The adult eyes of all these lines displayed a small-rough eye phenotype (eyes smaller than normal, with irregular, rough phenotype). This indicates that both lats and h-lats genes have the same biological effect when they are overexpressed in the developing Drosophila eye.

# 9.2. HUMAN H-LATS GENE CAN REPLACE THE DROSOPHILA HOMOLOG TO PREVENT DEATH IN DROSOPHILA ANIMALS HAVING MUTANT DROSOPHILA LATS

The Drosophila lats cDNA was cloned into the pCaSpeR-hs vector (Thummel and Pirrotta, 1992, Drosophila Inform. Service 71:150) for germ line transformation of

15 Drosophila. Three of the transformed lines were tested and were able to rescue the lethality of the latsal/latsal, latsal and latsal animals after one hour heat shock for every 24 hours during larval and pupal development. The human h-lats cDNA (in a XhoI (blunted)-XbaI fragment) from pBS(SK)-h-lats CDNA (Fig. 10) was cloned into the HpaI-XbaI sites of the pCaSpeR-hs vector, to produce plasmid pCaSpeR-hs-h-lats (Fig. 14). Plasmid pCaSpeR-hs-h-lats was used for germ line transformant. Three of the pCaSpeR-hs-h-lats transformant lines were tested and were able to rescue the lethality of the latsal and latsal animals under the same conditions used in rescuing experiments for the Drosophila gene.

# 10. HUMAN LATS EXPRESSION IS FOUND IN ALL NORMAL TISSUES TESTED AND IS ABSENT IN A LARGE NUMBER OF TUMOR CELL LINES

The expression of human lats RNA was investigated in various adult tissues. A 1.2 kb BamHI fragment of the h-lats cDNA was used as a <sup>32</sup>P-labeled probe for Northern analysis. Hybridization was to a nylon membrane containing polyA+ RNA from various human fetal and adult tissues, obtained from Clontech. The Northern analysis was carried



out according to the recommended instructions of the manufacturer (Clontech). The results are shown in Figure 15. h-lats was expressed in every tissue tested (fetal brain, fetal lung, fetal liver, fetal kidney, adult spleen, adult thymus, adult prostate, adult testis, adult ovary, adult small intestine, adult colon, and adult blood leukocytes). Expression was higher in fetal tissues than in adult tissues.

## 10.2. HUMAN LATS EXPRESSION IN VARIOUS TUMOR CELL LINES

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The <sup>32</sup>P-labeled BamHI fragment of h-lats was used as a probe for Northern analysis, for hybridization to total RNAs isolated from 42 different human tumor cell lines (obtained from the American Type Culture Collection, Rockville, MD). No h-lats expression was detected in 20 of the tumor lines (48%). The name and tissue origin of the tumor cell lines tested, and the results of the Northern analysis are presented in Table 3.

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	T	a	b	1	e	3
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	Name of tumor lines	Tumor Origin	Expression det	alyses
25			<u>YES</u>	<u>NO</u>
	5637	Bladder		<u>ио</u>
	RT4	Bladder	土*	
	HT-1376	Bladder		x
	HT-1197	Bladder		X
		B = = 4	v	
30	BT-20	Breast	X	
	BT-474	Breast	X	
	ZR-75-1	Breast		X
	ZR-75-30	Breast	X	
	BT-549	Breast		X
	MDA-MB-453	Breast		X
	MDA-MB-435S	Breast		X
	HBL-100	Breast		X
35	• •••	00100		х
	LoVo	Colon	v	7.
	HT-29	Colon	X	
	HCT116	Colon	X	
	LS 180	Colon		X
	DLD-1	Colon	X	
	WiDr	Colon	X	

V	VO 96/3		PCT/I	JS96/04101
	SW480	Colon	x	
	Caco-2	Colon	±	
	HEL 92.1.7	Erythroleukemia	x	
	MOLT-4	Leukemia	X	
	CEM-CM3	Leukemia	X	
5	K-562	Leukemia	· X	
	Jurkat	Leukemia		X
	HUT 78	Lymphoma	x	
	SK-LU-1	Lung	•	x
	A-427	Lung		X
	Calu-1	Lung	x	•
10	NCI-H69	Lung	x	
	SK-MEL-3	Melanoma		х
	SK-MEL-28	Melanoma		X
	SK-MEL-31	Melanoma		x
	MIA PaCa-2	Pancreas		x
	BxPC-3	Pancreas		×
15	Hs 700T	Pancreas	x	
	Hs 766T	Pancreas	x	A, ē
	RD	Sarcoma		x
	A-204	Sarcoma		×
	AN3 CA	Uterine	x	

SK-UT-1

HEC-1-A

Uterine

Uterine

X

<sup>\*:</sup> weak signal

Thus, 48% of the tumor cell lines tested had no detectable h-lats expression, whereas 100% of the normal tissues tested had detectable h-lats expression. It should be noted that the 48% figure may be an underestimate of the actual number of tumor cell lines that had decreased lats protein level or activity relative to normal tissue, since

while lack of lats RNA (i.e., a transcriptional block) allows the conclusion that no lats protein is made, tumor cells that expressed the lats RNA may still have had no or low lats protein levels and/or activity due to the possible existence

<sup>35</sup> of a translational block or the presence of mutation(s) in an expressed lats protein.

## 11. <u>DEPOSIT OF MICROORGANISM</u>

pBS(KS)-h-lats was deposited on March 24, 1995 with the American Type Culture Collection, 1201 Parklawn Drive,

- 5 Rockville, Maryland 20852, under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedures, and assigned Accession No. 69769.
- Described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing
- 15 description and accompanying figures. Such modifications are intended to fall within the scope of the appended claims.

Various references are cited herein, the disclosures of which are incorporated by reference in their entireties.

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## SEQUENCE LISTING

## (1) GENERAL INFORMATION:

- (i) APPLICANT: Xu, Tian Tao, Wufan Wang, Weiyi Zhang, Sheng Yu, Wan
- (ii) TITLE OF INVENTION: NUCLEOTIDE AND PROTEIN SEQUENCES OF LATS GENES AND METHODS BASED THEREON
- (iii) NUMBER OF SEQUENCES: 16
- (iv) CORRESPONDENCE ADDRESS:
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  - (B) STREET: 1155 Avenue of the Americas
  - (C) CITY: New York
  - (D) STATE: New York
  - (E) COUNTRY: USA
  - (F) ZIP: 10036-2711
- (V) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Floppy disk
  - (B) COMPUTER: IBM PC compatible
  - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
  - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: To Be Assigned
  - (B) FILING DATE: On Even Date Herewith
  - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Misrock, S. Leslie
  - (B) REGISTRATION NUMBER: 18,872
  - (C) REFERENCE/DOCKET NUMBER: 6523-007
  - (ix) TELECOMMUNICATION INFORMATION:
    - (A) TELEPHONE: (212) 790-9090
    - (B) TELEFAX: (212) 869-9741/8864
    - (C) TELEX: 66141 PENNIE
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 5720 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: double (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: cDNA
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS
    - (B) LOCATION: 1103..4402
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

ATCTAGCACG ACGGCAGCAA CAAAACCACG AATTAATTTT ACTAAATTTA AGCCAAACGC



<del>-</del>	
GCATCGGAAA TGCCTGAAAA TGCGATTGAA TGCACGCGAA AAGTGATGGG TTGCG	AACGC 120
GAGTGAATCA AGTGAAAATA CGTCGGCAAA TATCAGCGAA TTGTCGTCAA AAGGC	AAGGA 180
ANANCGGAGA ANANGAGGAA ANGCANTANG TGCCGTGTGT GGGAANCGCG ANANA	GGCGA 240
GAACAAAGAG GCGAAAAGCG AGGAAATTGC GTGGAAAAACG TGGAAAACGC GAAGA	agcga 300
AGCTCCAAGT TGGCCGCCAT CGATTCGTGC GTAGGATCAA TTAAGATTCC GAGTG	GTCGA 360
GAATCGGCTC AAATCAAATT AAAATCAACT AATATTTTGG TATTCAGATA TTCAA	ATGGA 420
ATTCATTCAT CGCCTGCGAC TTTTATTCGG ATCTGCCAAC TATTTTTGAA TTTGA	ATTGT 480
GTGTCTGCGG CTGGCGCAGA ATCTCTGATA AAGCAGAGGA ATAAAATCGG AAGAA	CAACA 540
ANTACAANTA CAANTGAANT GCGGGGAGCA GTATTTACAT GCCAANTGAA TGCTG	GATAG 600
GCGAAAGGGG GGGTTTCTCT TATAATGCAA ATGTGAATGT GAATGCGAAT GCGAA	TGCGA 660
GTGGAAGAAT TCCCGGCGCG AGTGATAAAT AATCCGACGA CAAACAAAGC AGAAG	CCTAC 720
ACCGCGAGAA AGAGCAGCGC AAACACAATT ATCTTTATTG AGAGCAACAA TATCA	AGATC 780
GAGATAATAA AGCATCCTAA AACCCGCGCC TTAGTTCGTT TTAGTCTCGC CACGG	SATATA 840
GATATTCARA GGCARARAGG TGGTGTCGGC ATCGCCAGAC ARACRAGTAR AGCAT	CTATT 900
TCATACAAAA CAACCAATTA AATAATAATA AAAATAATAA TAATCGTAGA GAGGC	AGAGC 960
CANATCHANT TCCCGGCCGC CGATGTGCCC CAGTGTGTGT GCGTGTGTGT GTGTG	TGTGC 1020
TGTGCTGTGC TGTGCGAGTG TTAGTGTGCG GAGCATTTCT GTGATATGAG TGCTA	AATGC 1080
CACAGGGCGA AGCAGCAGCA TC ATG CAT CCA GCG GGC GAA AAA AGG GGC Met His Pro Ala Gly Glu Lys Arg Gly	GGT 1132 Gly 10
CGC CCC AAT GAT AAA TAC ACG GCG GAA GCC CTC GAG AGC ATC AAG Arg Pro Asn Asp Lys Tyr Thr Ala Glu Ala Leu Glu Ser Ile Lys 15	CAG 1180 Gln
GAC CTA ACC CGA TTT GAA GTA CAA AAT AAC CAT AGG AAT AAT CAG Amp Leu Thr Arg Phe Glu Val Gln Am Am Him Arg Am Am Gln 30 35 40	AAT 1228 Asn
TAC ACA CCT CTG CGA TAC ACG GCG ACC AAC GGA CGC AAC GAT GCA Tyr Thr Pro Leu Arg Tyr Thr Ala Thr Asn Gly Arg Asn Asp Ala 45	CTT 1276 Leu
ACT CCT GAC TAT CAC CAC GCC AAG CAG CCG ATG GAG CCG CCA CCC Thr Pro Asp Tyr His His Ala Lys Gln Pro Met Glu Pro Pro Pro 60 65 70	TCC 1324 Ser
GCC TCT CCT GCT CCG GAC GTG GTC ATA CCG CCG CCC CCC GCC ATT	GTA 1372
Ala Ser Pro Ala Pro Asp Val Val Ile Pro Pro Pro Pro Ala Ile 75 80 85	90
GGT CAG CCC GGA GCC GGC TCC ATA TCC GTA TCC GGT GTG GGC GTT Gly Gln Pro Gly Ala Gly Ser Ile Ser Val Ser Gly Val Gly Val 95 100	GGA 1420 Gly
GTG GTG GGT GTG GCG AAC GGA CGT GTG CCA AAG ATG ATG ACG GCC Val Val Gly Val Ala Asn Gly Arg Val Pro Lys Met Met Thr Ala 110 115	CTA 1468 Leu



Met	Pro	Aøn 125	Lys	Leu	Ile	Arg	Lys 130	Pro	Ser	Ile	Glu	Arg 135	Asp	Thr	Ala	
														GCC	GGT Gly	1564
AGC Ser 155	TCC	CGA Arg	TCG Ser	GAC Asp	AGC Ser 160	CCC Pro	CAT His	TCG Ser	CAC His	CAC His 165	ACC Thr	CAC His	CAG Gln	CCG Pro	AGC Ser 170	1612
TCG Ser	<b>A</b> GG <b>A</b> rg	ACG Thr	GTG Val	GGT Gly 175	AAT Asn	CCA Pro	GGT Gly	GGA Gly	AAT Aen 180	GGT Gly	GGA Gly	TTT Phe	TCT Ser	CCG Pro 185	TCG Ser	1660
CCA Pro	AGC Ser	GGT Gly	TTC Phe 190	AGT Ser	GAG Glu	GTG Val	GCT Ala	CCA Pro 195	CCG Pro	GCG Ala	CCG Pro	CCG Pro	CCA Pro 200	CGC Arg	AAT Asn	1708
CCC Pro	ACC	GCC Ala 205	TCC Ser	AGC Ser	GCG Ala	GCC Ala	ACG Thr 210	CCC Pro	CCA Pro	CCG Pro	CCA Pro	GTG Val 215	CCG Pro	CCC Pro	ACC Thr	1756
AGC Ser	CAG Gln 220	GCG Ala	TAC Tyr	GTG Val	AAG Lys	CGG Arg 225	CGA Arg	TCA Ser	CCG Pro	GCC Ala	CTG Leu 230	AAC Asn	AAC Asn	CGC Arg	CCG Pro	1804
CCG Pro 235	GCG Ala	ATA Ile	GCG Ala	CCA Pro	CCC Pro 240	ACT Thr	CAG Gln	CGA Arg	GGC Gly	AAC Asn 245	TCA Ser	CCT Pro	GTA Val	ATA Ile	ACC Thr 250	1852
CAA Gln	AAC Asn	GGG Gly	CTG Leu	AAG Lys 255	AAC Asn	CCG Pro	CAG Gln	CAG Gln	CAG Gln 260	TTG Leu	ACG Thr	CAG Gln	CAG Gln	CTG Leu 265	AAG Lys	1900
TCC	CTG Leu	AAC Aen	CTA Leu 270	TAC Tyr	CCA Pro	GGC Gly	GGA Gly	GGC Gly 275	AGT Ser	GGA Gly	GCA Ala	GTG Val	GTG Val 280	GAG Glu	CCA Pro	1948
CCG Pro	CCG Pro	CCC Pro 285	TAC Tyr	CTA Leu	ATT Ile	CAA Gln	GGC Gly 290	GGA Gly	GCC Ala	GGA Gly	GGA Gly	GCA Ala 295	GCA Ala	CCG Pro	CCG Pro	1996
CCG Pro	CCA Pro 300	CCA Pro	CCC Pro	AGT Ser	TAC Tyr	ACG Thr 305	GCC Ala	TCC Ser	ATG Met	CAG Gln	TCG Ser 310	CGG Arg	CAG Gln	TCG Ser	CCC Pro	2044
ACA Thr 315	CAA Gln	TCC Ser	CAA Gln	CAA Gln	TCG Ser 320	GAC Asp	TAC Tyr	AGG Arg	AAA Lys	TCC Ser 325	CCG Pro	AGC Ser	AGT Ser	GGG Gly	ATA Ile 330	2092
TAC Tyr	TCG Ser	GCC Ala	ACC Thr	TCG Ser 335	GCG Ala	GGC Gly	TCG Ser	CCG Pro	AGC Ser 340	CCC Pro	ATA Ile	ACT Thr	GTG Val	TCG Ser 345	CTG Leu	2140
CCG Pro	CCG Pro	GCG Ala	CCG Pro 350	CTG Leu	GCG Ala	AAG Lyb	CCA Pro	CAA Gln 355	CCA Pro	CGA Arg	GTC Val	TAC Tyr	CAG Gln 360	GCC Ala	AGG Arg	2188
AGT Ser	CAG Gln	CAG Gln 365	CCG Pro	ATC Ile	ATC Ile	ATG Met	CAG Gln 370	AGT Ser	GTG Val	AAG Lyв	AGC Ser	ACG Thr 375	CAG Gln	GTC Val	CAA Gln	2236
AAG Lys	CCC Pro 380	GTG Val	CTG Leu	CAA Gln	ACA Thr	GCA Ala 385	GTG Val	GCG Ala	CGC Arg	CAA Gln	TCG Ser 390	CCA Pro	TCG Ser	AGT Ser	GCC Ala	2284

TCG Ser 395	GCC Ala	<b>A</b> GC Ser	AAT Asn	TCA Ser	CCA Pro 400	GTC Val	CAC His	GTG Val	Leu	GCC Ala 405	GCT Ala	CCA Pro	CCC Pro	TCT Ser	TAC Tyr 410	2332
CCT Pro	CAG Gln	AAG Lys	TCC Ser	GCG Ala 415	GCA Ala	GTG Val	GTG Val	CAG Gln	CAG Gln 420	CAG Gln	CAA Gln	CAG Gln	Ala	GCA Ala 425	GCG Ala	2380
GCG Ala	GCC Ala	CAC His	CAG Gln 430	CAG Gln	CAG Gln	CAT His	CAG Gln	CAC His 435	CAG Gln	CAA Gln	TCC Ser	AAA Lys	CCA Pro 440	CCA Pro	ACG Thr	2428
CCA Pro	ACC Thr	ACA Thr 445	CCG Pro	CCC Pro	TTG Leu	GTG Val	GGT Gly 450	CTG Leu	AAC Asn	AGC Ser	AAG Lys	CCC Pro 455	AAT Asn	TGC Cys	CTG Leu	2476
GAG Glu	CCA Pro 460	ccg Pro	TCC Ser	TAT Tyr	GCC Ala	AAG Lys 465	AGC Ser	ATG Met	CAG Gln	GCC Ala	AAG Lys 470	GCG Ala	GCC Ala	ACG Thr	GTG Val	2524
GTA Val 475	CAG Gln	CAG Gln	CAG Gln	CAA Gln	CAG Gln 480	CAG Gln	CAG Gln	CAA Gln	CAA Gln	CAG Gln 485	CAG Gln	GTC Val	CAG Gln	CAG Gln	CAG Gln 490	2572
CAG Gln	GTG Val	CAA Gln	CAG Gln	CAG Gln 495	CAG Gln	CAA Gln	CAG Gln	CAG Gln	CAA Gln 500	CAG Gln	CAA Gln	CTG Leu	CAG Gln	GCC Ala 505	TTG Leu	2620
AGG Arg	GTG Val	CTC	CAG Gln 510	Ala	CAG Gln	GCT Ala	CAG Gln	AGG Arg 515	Glu	CGG	GAT Asp	CAA Gln	CGG Arg 520	GAG Glu	CGG Arg	2668
G <b>AA</b> Glu	CGG Arg	GAT Asp 525	Gln	CAG Gln	AAG Lys	CTG	GCC Ala 530	Asn	GGA Gly	AAT Asn	CCT	GGC Gly 535	CGG Arg	CAG Gln	ATG Met	2716
CTT Leu	CCG Pro 540	Pro	ccc Pro	ccc Pro	TAT Tyr	Gln 545	Ser	AAC	AAC ABD	AAC	AAC ABD 550	ABD	AGC Ser	GAG Glu	ATC	2764
<b>AAA</b> Lys 555	Pro	CCG Pro	AGC Ser	TGC Cys	AAC ABD 560	yeu	AAC ABr	AAC ABr	ATA	CAG Gln 565	Ile	AGC Ser	AAC	AGC Ser	AAC Asn 570	2812
CTG Leu	GCG	ACC Thi	ACI Thi	CC) Pro 575	Pro	ATI Ile	CCC Pro	CCI Pro	GCC Ala 580	Lys	TAC Tyl	AAT Asn	AAC Asn	AAC ABr 585	TCC Ser	2860
TCC Ser	AAC Aar	ACC Thi	G GG( r Gl) 59(	y Ala	AAT ABI	AGC Sei	TCC Sei	GG( Gl) 599	λ GT?	Ser	AAC ABI	GGA n Gly	Ser 600	T 111	GGC Gly	2908
ACC Thr	ACC Thi	GC Al	a Se	C TCC r Se	TCC r Sei	ACC Thi	Ser 610	r Cyi	C AAC B Lys	AAC Lys	S ATO	C AAG E Lys 619	HIE	C GCC B Ala	C TCG A Ser	2956
CCC	2 ATC	Pr	G GA	G CG	C AAG g Ly	S AAG B Ly: 62!	B Ile	C TC e Se:	C AAG	G GAG	G AAG Ly: 63	B GT	G GAG	G GAG	G CGC	3004
AAG Lys 635	Gl	G TT u Ph	c cg e Ar	C AT	C AGG e Arg 640	g G1:	G TA	C TO	G CC	G CA G G1: 64:	U AT	C TTO	C AAG E Lys	G TTO	C TTC e Phe 650	
ATC Met	G GA	G CA u Gl	G CA n Hi	C AT 8 Il 65	e Gl	G AA u Ab	C GT n Va	G AT 1 11	C AA e Ly: 66	B Se	G TA r Ty	T CG	C CAG	G CG n Ar 66	C ACG g Thr 5	3100



					CTG Leu											3148
					GAG Glu											3196
					AAG Lys											3244
					GGA Gly 720											3292
					TCG Ser											3340
					AAG Lys											3388
					GAA Glu											3436
TAC	AGC Ser 780	TTC Phe	CAG Gln	GAC Asp	AAG Lys	GAT Asp 785	AAT ABN	CTG Leu	TAC Tyr	TTT Phe	GTG Val 790	ATG Met	GAC Asp	TAC Tyr	ATA Ile	3484
					ATG Met 800											3532
GAG Glu	GAA Glu	CTG Leu	GCC Ala	AGA Arg 815	TTC Phe	TAC Tyr	ATC Ile	GCC Ala	GAG Glu 820	GTC Val	ACC Thr	TGC Cys	GCC Ala	GTG Val 825	GAC Asp	3580
AGC Ser	GTT Val	CAC His	AAA Lys 830	ATG Met	GLY	TTC Phe	ATT Ile	CAC His 835	AGA Arg	GAC Asp	ATC Ile	AAG Lys	CCT Pro 840	GAC Asp	AAC Asn	3628
ATA Ile	CTC Leu	ATC Ile 845	GAT Asp	AGG Arg	GAC Asp	GGA Gly	CAC His 850	ATA Ile	AAG Lys	CTC Leu	ACC Thr	GAC Asp 855	TTT Phe	GGC Gly	CTG Leu	3676
TGC Cys	ACG Thr 860	GGA Gly	TTC Phe	CGA Arg	TGG Trp	ACG Thr 865	CAC His	AAC Asn	TCG Ser	AAG Lys	TAC Tyr 870	TAC Tyr	CAG Gln	GAG Glu	AAC Asn	3724
GGC Gly 875	AAT Abn	CAC His	TCG Ser	CGC Arg	CAG Gln 880	GAC Asp	TCG Ser	ATG Met	GAG Glu	CCC Pro 885	TGG Trp	GAG Glu	GAA Glu	TAC Tyr	TCC Ser 890	3772
GAG Glu	AAC Asn	GGA Gly	CCG Pro	AAG Lys 895	CCC Pro	ACC Thr	GTG Val	CTG Leu	GAG Glu 900	AGG Arg	CGA Arg	CGG Arg	ATG Met	CGC Arg 905	GAT Asp	3820
CAC His	CAA Gln	AGA Arg	GTC Val 910	CTG Leu	GCC Ala	CAC His	TCG Ser	CTG Leu 915	GTG Val	GGC Gly	ACC Thr	CCG Pro	AAC Asn 920	TAC Tyr	ATA Ile	3868
GCT Ala	CCC Pro	GAG Glu 925	GTG Val	CTG Leu	GAG Glu	AGG Arg	AGT Ser 930	GGG Gly	TAC Tyr	ACG Thr	CAG Gln	CTG Leu 935	TGC Cys	GAC Asp	TAC Tyr	3916

TGG AGC GTG GGC GTC ATC CTT TAC GAG ATG CTG GTG GGT CAG CCG CCC Trp Ser Val Gly Val Ile Leu Tyr Glu Met Leu Val Gly Gln Pro Pro 940 945	3964
TTT CTG GCC AAC AGT CCG CTG GAA ACG CAA CAA AAG GTC ATC AAC TGG Phe Leu Ala Asn Ser Pro Leu Glu Thr Gln Gln Lys Val Ile Asn Trp 955 960 965 970	4012
GAG AAA ACG CTG CAT ATT CCG CCG CAG GCC GAG TTA TCC CGC GAG GCT Glu Lys Thr Leu His Ile Pro Pro Gln Ala Glu Leu Ser Arg Glu Ala 975 980 985	4060
ACG GAC TTG ATA AGG AGG CTC TGT GCG TCG GCT GAC AAG CGG CTG GGC Thr Asp Leu Ile Arg Arg Leu Cys Ala Ser Ala Asp Lys Arg Leu Gly 990 995	4108
AAG AGC GTG GAC GAG GTC AAG AGC CAC GAC TTC TTC AAG GGC ATC GAC Lys Ser Val Asp Glu Val Lys Ser His Asp Phe Phe Lys Gly Ile Asp 1005 1010 1015	4156
TTT GCG GAC ATG CGG AAG CAG AAA GCG CCC TAC ATA CCG GAA ATC AAG Phe Ala Asp Met Arg Lys Gln Lys Ala Pro Tyr Ile Pro Glu Ile Lys 1020 1025. 1030	4204
CAC CCA ACG GAC ACA TCC AAC TTT GAT CCC GTG GAT CCG GAG AAG CTG His Pro Thr Asp Thr Ser Asn Phe Asp Pro Val Asp Pro Glu Lys Leu 1035 1040 1045 1050	4252
CGC TCG AAT GAC TCC ACC ATG AGC AGC GGC GAT GAT GTC GAC CAG AAT Arg Ser Asn Asp Ser Thr Het Ser Ser Gly Asp Asp Val Asp Gln Asn 1055 1060 1065	4300
GAC CGC ACT TTC CAC GGC TTT TTC GAA TTT ACC TTC CGT CGC TTC TTC Asp Arg Thr Phe His Gly Phe Phe Glu Phe Thr Phe Arg Arg Phe Phe 1070 1080	4348
GAC GAC AAG CAG CCG CCG GAT ATG ACG GAC GAT CAG GCG CCG GTT TAC Asp Asp Lys Gln Pro Pro Asp Met Thr Asp Asp Gln Ala Pro Val Tyr 1085 1090 1095	4396
GTC TGA AATGGATGCT CTCCATGTGC CCAACACCAA CACCCCGCCC CCGAATCATT Val * 1100	4452
GTTAGTCAAA TAGTCACAAA AAGGGGATAG AAACCATTGA GTGGGCTTGC ATTGTAAAGG	4512
AAGCGTGGCT ATAGAATGAA ACTATCTATA TACATTATAT AAATTATAGG AGACAGTAGA	4572
GGCGGGAGCT ACGTATATAC ATACAAATAA TATACATATA TTTGATATAT ATATATATAT	4632
ATATGCCGTA GGGCATGAAC TGAATAAATA TAAAACGGAG CCGAGTAGAG ATGAAACGAG	4692
AGGAGCGAGT CAGGACCTTC GACCTTTAAC TGAACATAGT ATATCCTTGT GCACTACTAC	4752
TCCACAACAA ATATATATT TTAAATTGTT AGAATTCAAA AGGGACCAAC TGGAAATCGA	4812
ACCTTTCTGG TGCTCAAAGC AAAGCAAAGC AAAGCAAAAC AAAACGCCTT AAACTAAATG	4872
AGACGCGAAT TTACCCAACC ACTTCACTCC TCTCCTTTCT CCACCTCCGA TCGGTGGCCG	4932
GATTCGAACT CAGCAGGCTG GTTGCATCCG GCCATCCCAT TGACTTCCCA TTCAGAATTG	4992
AGATTGCGAG GTGTGCGATG GAGAACGAAC GGAGACCAAA AGTCGCACGG CAGCGATATA	5052
AGCGGGTCTT ATAAGCCTAA TCTAAATCTA AACTGGGAGA ACAGGACCTA TGTATGTCCT	5112
GCTATCCAAT TCGTCTATCA CTGCTCTTCA TCTGTGTACG ACCCCCACCC CCCCCCCCC	5172



					101/03	0/0410)
Catccaaaag	AACAAACTTA	GACGTAGCCT	ATGTGAAAAG	CTAGCAATGT	TAGACCAACT	5232
TGTTGAATGC	CAAATGAAAT	TGTTTAGCCC	CACGAGGAAA	ACGCGGGGGA	AATTCAACAC	5292
TTATTCTCTG	ATAGCAAACG	GAAAAGAAAG	AAAGAAAAA	AAAAACAGAA	ACAGTACGAG	5352
<b>AAAATT</b> GTAA	TCTTCTTAAT	GTAATATTGT	AAAGAACACG	TTAATTGTAA	TCTATGCTAG	5412
agitgigiag	CGCCCTAAGA	TGTTTTTAG	TTTATAGACC	GCTAACCGTA	ATCTAGTTTA	5472
attectaaca	CTAAGCGAGA	GTACAGTACA	TTGGTTTTTT	TGTTTGTCGT	AGGTTCGTTG	5532
GAAAATGCTT	AACGGGAAAC	GATTTGTTTT	TCTCTTTAAT	TAGCTTCAGT	TTGTATGTGC	5592
GTGTGTTTT	ATTATGACTT	ATATATAGTC	CATCTGAATA	TTCGTGGATG	GAGCCTATTT	5652
<b>TAAAT</b> GTGAG	ATCGAGCTAA	TTGAAGGAAA	TACAAACAAA	CTCTGTGTGC	CTTGGCCAAT	5712
TAGTTTAC						5720
(2) INFORM	ATION FOR SE	Q ID NO:2:				
· (i)	SEQUENCE CH	IARACTERISTI				

- (B) TYPE: amino acid (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met His Pro Ala Gly Glu Lys Arg Gly Gly Arg Pro Asn Asp Lys Tyr

1 10 15

Thr Ala Glu Ala Leu Glu Ser Ile Lys Gln Asp Leu Thr Arg Phe Glu 20 25 30

Val Gln Asn Asn His Arg Asn Asn Gln Asn Tyr Thr Pro Leu Arg Tyr

Thr Ala Thr Asn Gly Arg Asn Asp Ala Leu Thr Pro Asp Tyr His His

Ala Lys Gln Pro Het Glu Pro Pro Pro Ser Ala Ser Pro Ala Pro Asp 65

Val Val Ile Pro Pro Pro Pro Ala Ile Val Gly Gln Pro Gly Ala Gly

Ser Ile Ser Val Ser Gly Val Gly Val Gly Val Gly Val Ala Asn 105

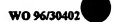
Gly Arg Val Pro Lys Met Met Thr Ala Leu Met Pro Asn Lys Leu Ile 120

Arg Lys Pro Ser Ile Glu Arg Asp Thr Ala Ser Ser His Tyr Leu Arg

Cys Ser Pro Ala Leu Asp Ser Gly Ala Gly Ser Ser Arg Ser Asp Ser

Pro His Ser His His Thr His Gln Pro Ser Ser Arg Thr Val Gly Asn 170

Pro Gly Gly Asn Gly Gly Phe Ser Pro Ser Gly Phe Ser Glu 185



Val Ala Pro Pro Ala Pro Pro Pro Arg Asn Pro Thr Ala Ser Ser Ala 200 Ala Thr Pro Pro Pro Pro Val Pro Pro Thr Ser Gln Ala Tyr Val Lys Arg Arg Ser Pro Ala Leu Asn Asn Arg Pro Pro Ala Ile Ala Pro Pro Thr Gln Arg Gly Asn Ser Pro Val Ile Thr Gln Asn Gly Leu Lys Asn 250 Pro Gln Gln Gln Leu Thr Gln Gln Leu Lys Ser Leu Asn Leu Tyr Pro Gly Gly Gly Ser Gly Ala Val Val Glu Pro Pro Pro Pro Tyr Leu Ile 280 Gln Gly Gly Ala Gly Gly Ala Ala Pro Pro Pro Pro Pro Pro Ser Tyr Thr Ala Ser Het Gln Ser Arg Gln Ser Pro Thr Gln Ser Gln Gln Ser Amp Tyr Arg Lys Ser Pro Ser Ser Gly Ile Tyr Ser Ala Thr Ser Ala Gly Ser Pro Ser Pro Ile Thr Val Ser Leu Pro Pro Ala Pro Leu Ala 345 Lys Pro Gln Pro Arg Val Tyr Gln Ala Arg Ser Gln Gln Pro Ile Ile Met Gln Ser Val Lys Ser Thr Gln Val Gln Lys Pro Val Leu Gln Thr Ala Val Ala Arg Gln Ser Pro Ser Ser Ala Ser Ala Ser Asn Ser Pro Val His Val Leu Ala Ala Pro Pro Ser Tyr Pro Gln Lys Ser Ala Ala 405 Val Val Gln Gln Gln Gln Ala Ala Ala Ala His Gln Gln Gln 425 His Gln His Gln Gln Ser Lys Pro Pro Thr Pro Thr Pro Pro Leu Val Gly Leu Asn Ser Lys Pro Asn Cys Leu Glu Pro Pro Ser Tyr Ala Lys Ser Met Gln Ala Lys Ala Ala Thr Val Val Gln Gln Gln Gln Gln Gin Gin Gin Gin Gin Val Gin Gin Gin Val Gin Gin Gin Gin 490 Gin Gin Gin Gin Gin Leu Gin Ala Leu Arg Val Leu Gin Ala Gin Ala Gln Arg Glu Arg Asp Gln Arg Glu Arg Glu Arg Asp Gln Gln Lys Leu Ala Asn Gly Asn Pro Gly Arg Gln Het Leu Pro Pro Pro Pro Tyr Gln Ser Asn Asn Asn Asn Ser Glu Ile Lys Pro Pro Ser Cys Asn

545 550 555 Asn Asn Asn Ile Gln Ile Ser Asn Ser Asn Leu Ala Thr Thr Pro Pro 565 570 Ile Pro Pro Ala Lys Tyr Asn Asn Asn Ser Ser Asn Thr Gly Ala Asn 585 Ser Ser Gly Gly Ser Asn Gly Ser Thr Gly Thr Thr Ala Ser Ser Ser Thr Ser Cys Lys Lys Ile Lys His Ala Ser Pro Ile Pro Glu Arg Lys Lys Ile Ser Lys Glu Lys Glu Glu Glu Arg Lys Glu Phe Arg Ile Arg Gin Tyr Ser Pro Gin Ala Phe Lys Phe Phe Met Glu Gin His Ile Glu Asn Val Ile Lys Ser Tyr Arg Gin Arg Thr Tyr Arg Lys Asn Gin Leu Glu Lys Glu Met His Lys Val Gly Leu Pro Asp Gln Thr Gln Ile Glu 680 Met Arg Lys Met Leu Asn Gln Lys Glu Ser Asn Tyr Ile Arg Leu Lys Arg Ala Lys Met Asp Lys Ser Met Phe Val Lys Leu Lys Pro Ile Gly 705 710 715 720 Val Gly Ala Phe Gly Glu Val Thr Leu Val Ser Lys Ile Asp Thr Ser Asn His Leu Tyr Ala Het Lys Thr Leu Arg Lys Ala Asp Val Leu Lys 745 Arg Asn Gln Val Ala His Val Lys Ala Glu Arg Asp Ile Leu Ala Glu Ala Asp Asn Asn Trp Val Val Lys Leu Tyr Tyr Ser Phe Gln Asp Lys Asp Asn Leu Tyr Phe Val Het Asp Tyr Ile Pro Gly Gly Asp Leu Het Ser Leu Leu Ile Lys Leu Gly Ile Phe Glu Glu Glu Leu Ala Arg Phe Tyr Ile Ala Glu Val Thr Cys Ala Val Asp Ser Val His Lys Met Gly Phe Ile His Arg Asp Ile Lys Pro Asp Asn Ile Leu Ile Asp Arg Asp Gly His Ile Lys Leu Thr Asp Phe Gly Leu Cys Thr Gly Phe Arg Trp Thr His Asn Ser Lys Tyr Tyr Gln Glu Asn Gly Asn His Ser Arg Gln Asp Ser Met Glu Pro Trp Glu Glu Tyr Ser Glu Asn Gly Pro Lys Pro Thr Val Leu Glu Arg Arg Arg Met Arg Asp His Gln Arg Val Leu Ala 905

284



WO 96/30402					101/03/004101
His Ser Leu Val	Gly Thr Pro	Asn Tyr 920	Ile Ala Pro	Glu Val Leu 925	Glu
Arg Ser Gly Tyr 930	Thr Gln Leu 935	Сув Авр	Tyr Trp Ser 940	Val Gly Val	Ile
Leu Tyr Glu Met 945	Leu Val Gly 950	Gln Pro	Pro Phe Leu 955	Ala Asn Ser	Pro 960
Leu Glu Thr Gln	Gln Lys Val 965	Ile Asn	Trp Glu Lys 970	Thr Leu His 975	Ile
Pro Pro Gln Ala 980	)	703			
Leu Cys Ala Ser 995	Ala Asp Lys	Arg Leu 1000	Gly Lys Ser	r Val Asp Glu 1005	Val
Lys Ser His Asp 1010	101	.5	20		
Gln Lys Ala Pro 1025	o Tyr Ile Pro 1030	Glu Ile	Lys His Pr 1035	o Thr Asp Thi	ser 1040
Asn Phe Asp Pr	o Val Asp Pro 1045	Glu Lys	Leu Arg Se 1050	r Asn Asp Ser 10	Thr 55
Met Ser Ser Gl	y Asp Asp Va	l Asp Gln 106	Asn Asp Ar	g Thr Phe Hi 1070	B Gly
Phe Phe Glu Ph 1075	e Thr Phe Ar	g Arg Phe 1080	Phe Asp As	p Lys Gln Pro 1085	o Pro
Asp Met Thr As	ip Asp Gln Al 10	a Pro Val 95	Tyr Val 1	100	
(2) INFORMATIO	ON FOR SEQ ID	ю:3:			
(A) (B) (C) (D)	ENCE CHARACTE LENGTH: 3984 TYPE: nuclei STRANDEDNESS TOPOLOGY: ur CULE TYPE: CL	base par c acid c double nknown			
(11) HOLE	CODE III.				
(ix) FEAT (A) (B)	URE: NAME/KEY: CI LOCATION: 2:	DS 313623			
(xi) SEQU	ENCE DESCRIP	TION: SEC	ID NO:3:		гсстаасс 60
ACCTTTGGGT TO	CTGGGACG GAC	TCTGGCC G	CCTCAGCGT (	CCCCCTCAG GC	CC010000
GCTGTCCAGG AG	CTCTGCTC TCC	CCTCCAG A	GTTAATTAT 1	TATATTGTA AA	grant and
ACAGTCCTGG GG					gCICIOOI.
TCTATCAAAT AA	AAGAAGTCC TTC	CTGTGGG (	CTACATATAT 1	AGATGTTTTU AT Me	t Lyb 1

AGG AGT GAA AAG CCA GAA GGA TAT AGA CAA ATG AGG CCT AAG ACC TTT Arg Ser Glu Lys Pro Glu Gly Tyr Arg Gln Met Arg Pro Lys Thr Phe 10



CCT Pro	GCC Ala 20	Ser	AAC	TAT Tyr	ACT	GTC Val 25	AGT Ser	AGC Ser	CGG Arg	CAA Gln	ATG Met 30	Leu	CAA Gln	GAA Glu	ATT	332
CGG Arg 35	Glu	TCC Ser	CTT Leu	AGG Arg	AAT Asn 40	Leu	TCT Ser	AAA Lys	CCA Pro	TCT Ser 45	GAT Asp	GCT Ala	GCT Ala	AAG Lys	GCT Ala 50	380
GAG Glu	CAT	AAC Asn	ATG Het	AGT Ser 55	Lys	ATG Het	TCA Ser	ACC	GAA Glu 60	Asp	CCT Pro	CGA Arg	CAA Gln	GTC Val 65	AGA Arg	428
AAT	CCA Pro	CCC Pro	AAA Lys 70	TTT	GGG Gly	ACG Thr	CAT His	CAT His 75	AAA Lys	GCC Ala	TTG Leu	CAG Gln	GAA Glu 80	ATT Ile	CGA Arg	476
AAC Asn	TCT	CTG Leu 85	CTT	CCA Pro	TTT Phe	GCA Ala	AAT Asn 90	GAA Glu	ACA Thr	AAT	TCT Ser	TCT Ser 95	CGG Arg	AGT Ser	ACT Thr	524
TCA Ser	GAA Glu 100	GTT Val	AAT Asn	CCA Pro	CAA Gln	ATG Met 105	CTT Leu	CAA Gln	GAC Asp	TTG Leu	CAA Gln 110	GCT Ala	GCT Ala	GGA Gly	TTT Phe	572
GAT Asp 115	GAG Glu	GAT Asp	ATG Met	GTT Val	ATA Ile 120	CAA Gln	GCT Ala	CTT Leu	CAG Gln	AAA Lys 125	ACT Thr	AAC Asn	AAC	AGA Arg	AGT Ser 130	620
ATA Ile	GAA Glu	GCA Ala	GCA Ala	ATT Ile 135	GAA Glu	TTC Phe	ATT Ile	AGT Ser	AAA Lys 140	ATG Met	AGT Ser	TAC Tyr	CAA Gln	GAT Asp 145	CCT Pro	668
CGA Arg	CGA Arg	GAG Glu	CAG Gln 150	ATG Met	GCT Ala	GCA Ala	GCA Ala	GCT Ala 155	GCC Ala	AGA Arg	CCT Pro	ATT	AAT Asn 160	GCC Ala	AGC Ser	716
ATG Met	AAA Lys.	CCA Pro 165	GGG Gly	TAA naA	GTG Val	CAG Gln	CAA Gln 170	TCA Ser	GTT Val	AAC Asn	CGC Arg	AAA Lys 175	CAG Gln	AGC Ser	TGG Trp	764
AAA Lys	GGT Gly 180	TCT Ser	AAA Lys	GAA Glu	TCC Ser	TTA Leu 185	GTT Val	CCT Pro	CAG Gln	AGG Arg	CAT His 190	GGC Gly	CCG Pro	CCA Pro	CTA Leu	812
GGA Gly 195	GAA Glu	AGT Ser	GTG Val	GCC Ala	TAT Tyr 200	CAT His	TCT Ser	GAG Glu	AGT Ser	CCC Pro 205	AAC Asn	TCA Ser	CAG Gln	ACA Thr	GAT Asp 210	860
GTA Val	GGA Gly	AGA Arg	CCT Pro	TTG Leu 215	TCT Ser	GGA Gly	TCT Ser	GGT Gly	ATA Ile 220	TCA Ser	GCA Ala	TTT Phe	GTT Val	CAA Gln 225	GCT Ala	908
CAC His	CCT Pro	AGC Ser	AAC Asn 230	GGA Gly	CAG Gln	AGA Arg	GTG Val	AAC Asn 235	CCC Pro	CCA Pro	CCA Pro	CCA Pro	CCT Pro 240	CAA Gln	GTA Val	956
AGG Arg	AGT Ser	GTT Val 245	ACT Thr	CCT Pro	CCA Pro	CCA Pro	CCT Pro 250	CCA Pro	AGA Arg	GGC Gly	CAG Gln	ACT Thr 255	CCC Pro	CCT Pro	CCA Pro	1004
AGA Arg	GGT Gly 260	ACA Thr	ACT Thr	CCA Pro	CCT Pro	CCC Pro 265	CCT Pro	TCA Ser	TGG Trp	GAA Glu	CCA Pro 270	AAC Asn	TCT Ser	CAA Gln	ACA Thr	1052
AAG Lys 275	CGC Arg	TAT Tyr	TCT Ser	GGA Gly	AAC Asn 280	ATG Met	GAA Glu	TAC Tyr	Val	ATC Ile 285	TCC Ser	CGA Arg	ATC Ile	TCT Ser	CCT Pro 290	1100

W U 90/30402					
GTC CCA CCT Val Pro Pro	GGG GCA TGG Gly Ala Trp 295	CAA GAG GGC Gln Glu Gly	TAT CCT CC Tyr Pro Pr 300	A CCA CCT CTC o Pro Pro Leu 305	AAC 1148 Asn
ACT TCC CCC Thr Ser Pro	ATG AAT CCT Met Asn Pro 310	CCT AAT CAA Pro Asn Glr 315	deta eru we	GA GGC ATT AGT GG Gly Ile Ser 320	TCT 1196 Ser
GTT CCT GTT Val Pro Val 325	GGC AGA CAA Gly Arg Gln	CCA ATC ATC Pro Ile Ile 330	C ATG CAG AG B Met Gln Se	er Ser Ser Lys 335	TTT 1244 Phe
AAC TTT CCA Asn Phe Pro 340	TCA GGG AGA Ser Gly Arg	CCT GGA ATO Pro Gly Met 345	C GIH VOH G	GT ACT GGA CAA ly Thr Gly Gln 50	ACT 1292 Thr
GAT TTC ATG Asp Phe Het 355	ATA CAC CAA Ile His Gln 360	WRU AGT AG	C CCT GCT GC 1 Pro Ala G 365	GC ACT GTG AAT ly Thr Val Asn	CGG 1340 Arg 370
CAG CCA CCA Gln Pro Pro	CCT CCA TAT	CCT CTG AC	A GCA GCT A Ir Ala Ala A 380	AT GGA CAA AGC sn Gly Gln Ser 385	CCT 1388
TCT GCT TTA Ser Ala Leu	CAA ACA GGG Gln Thr Gly 390	GGA TCT GC Gly Ser Al	a MIA PIO 3	CG TCA TAT ACA er Ser Tyr Thr 400	AAT 1436 Asn
GGA AGT ATT Gly Ser Ile 405	Pro Gln Se	T ATG ATG GT Het Met Va 410	rg cca aac a al Pro asn a	AGA AAT AGT CAT Arg Asn Ser His 415	r AAC 1484 s Asn
ATG GAA CTA Met Glu Leu 420	A TAT AAC AT 1 Tyr Asn Il	r AGT GTA CO e Ser Val Po 425	to ary men .	CAA ACA AAT TGG 31n Thr Asn Tr 130	G CCT 1532 p Pro
CAG TCA TC	r TCT GCT CC r Ser Ala Pr 44	O WIS GID S	CA TCC CCG 1 er Ser Pro 1 445	AGC AGT GGG CA Ser Ser Gly Hi	450
Ile Pro Th	r Trp Gln Pr 455	O WRU IIE I	460	TCA AAT TCT TT Ser ABn Ser Ph 46	5
Asn Pro Le	u Gly Asn Ai 470	g Ala Sel A	75	AAT TCT CAG CC Asn Ser Gln Pr 480	
Ala Thr Th	or Val Thr A. 35	490	TO ALE TIO	ATT CAA CAG CO Ile Gln Gln Pr 495	
Lys Ser Me 500	et Arg Val L	505	ju bed oin	ACT GCT TTA GC Thr Ala Leu A 510	
ACA CAC CO Thr His Po 515	ro Ser Trp I	TA CCA CAG ( le Pro Gln ) 20	CCA ATT CAA Pro Ile Gln 525	ACT GTT CAA CO	530
Pro Phe P	ro Glu Gly <sup>1</sup> 535	UL WIT SEL	540	GTG ATG CCA C Val Met Pro P	45
GCT GAA G Ala Glu A	CT CCA AAC 1 la Pro Asn 1 550	At GIH GT	CCA CCA CCA Pro Pro Pro 555	CCC TAC CCA A Pro Tyr Pro L 560	AA CAT 1916 ys His



CTG	CTC	CAC	CAA	AAC	CCA	de Carte	CTT	CCT	CCA	Th.C	CAC	TO B	) TO	AGT	226	1964
														Ser		1704
														AGT Ser		2012
														CAG Gln		2060
														GAG Glu 625		2108
														TTC Phe		2156
														CGT Arg		2204
														TTA Leu		2252
														GAA Glu		2300
AAT Asn	TAC Tyr	ATC Ile	CGT Arg	CTT Leu 695	AAA Lys	AGG Arg	GCT Ala	AAA Lys	ATG Met 700	GAC Asp	AAG Lys	TCT Ser	ATG Met	TTT Phe 705	GTG Val	2348
AAG Lys	ATA Ile	AAG Lys	ACA Thr 710	CTA Leu	GGA Gly	ATA Ile	GGA Gly	GCA Ala 715	TTT Phe	GGT Gly	GAA Glu	GTC Val	TGT Cys 720	CTA Leu	GCA Ala	2396
AGA Arg	AAA Lys	GTA Val 725	GAT Asp	ACT Thr	AAG Lys	GCT Ala	TTG Leu 730	TAT Tyr	GCA Ala	ACA Thr	AAA Lys	ACT Thr 735	CTT Leu	CGA Arg	AAG Lys	2444
AAA Lys	GAT Asp 740	GTT Val	CTT Leu	CTT Leu	CGA	AAT Asn 745	CAA Gln	GTC Val	GCT Ala	CAT His	GTT Val 750	AAG Lys	GCT Ala	GAG Glu	AGA Arg	2492
GAT Asp 755	ATC Ile	CTG Leu	GCT Ala	GAA Glu	GCT Ala 760	GAC Asp	AAT Asn	GAA Glu	TGG Trp	GTA Val 765	GTT Val	CGT Arg	CTA Leu	TAT Tyr	TAT Tyr 770	2540
TCA Ser	TTC Phe	CAA Gln	GAT Asp	AAG Lys 775	GAC Asp	AAT Asn	TTA Leu	TAC Tyr	TTT Phe 780	GTA Val	ATG Met	GAC Asp	TAC Tyr	ATT Ile 785	CCT Pro	2588
GGG Gly	GGT Gly	GAT Asp	ATG Met 790	ATG Met	AGC Ser	CTA Leu	TTA Leu	ATT Ile 795	AGA Arg	ATG Met	GGC Gly	ATC Ile	TTT Phe 800	CCA Pro	GAA Glu	2636
AGT Ser	CTG Leu	GCA Ala 805	CGA Arg	TTC Phe	TAC Tyr	ATA Ile	GCA Ala 810	GAA Glu	CTT Leu	ACC Thr	TGT Cys	GCA Ala 815	GTT Val	GAA Glu	AGT Ser	2684
GTT Val	CAT His 820	AAA Lys	ATG Met	GGT Gly	TTT Phe	ATT Ile 825	CAT His	AGA Arg	GAT Asp	ATT Ile	AAA Lys 830	CCT Pro	GAT Asp	AAT Asn	ATT Ile	2732

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TTG Leu 835	ATT Ile	GAT Asp	CGT Arg	GAT Asp	GGT Gly 840	CAT His	ATT Ile	AAA Lys	Leu	ACT Thr 845	GAC Asp	TTT Phe	GGC Gly	CTC Leu	TGC Cys 850	2780
ACT Thr	GCC Gly	TTC Phe	AGA Arg	TGG Trp 855	ACA Thr	CAC Hib	GAT Asp	TCT Ser	AAG Lys 860	TAC Tyr	TAT Tyr	CAG Gln	AGT Ser	GGT Gly 865	GAC Asp	2828
CAT His	CCA Pro	CGG Arg	CAA Gln 870	GAT Asp	AGC Ser	ATG Met	GAT Asp	TTC Phe 875	Ser	AAT Asn	GAA Glu	TGG Trp	GGG Gly 880	GAT Asp	CCC Pro	2876
TCA Ser	AGC Ser	TGT Cys 885	CGA Arg	TGT Cys	GGA Gly	GAC Asp	AGA Arg 890	CTG Leu	AAG Lys	CCA Pro	TTA Leu	GAG Glu 895	CGG Arg	AGA Arg	GCT Ala	2924
GCA Ala	CGC Arg 900	CAG Gln	CAC His	CAG Gln	CGA Arg	TGT Cys 905	CTA Leu	GCA Ala	CAT His	TCT Ser	TTG Leu 910	GTT Val	GGG Gly	ACT	CCC Pro	2972
AAT Asn 915	Tyr	ATT	GCA Ala	CCT Pro	GAA Glu 920	Val	TTG Leu	CTA Leu	CGA Arg	ACA Thr 925	GGA Gly	TAC Tyr	ACA Thr	CAG Gln	TTG Leu 930	3020
TGT Cys	GAT Asp	TGG Trp	TGG	AGT Ser 935	Val	GGT	GTT Val	ATT	CTT Leu 940	Phe	GAA Glu	ATG Met	TTG Leu	GTG Val 945	GGA	3068
CAA Gln	CCT Pro	CCT Pro	TTC Phe 950	Lev	GCA Ala	CAA Gln	ACA Thr	Pro 955	Leu	GAA Glu	ACA Thr	CAP Glr	ATG Met 960	ry	GTT Val	3116
ATC Ile	AAC ABD	TGG Trp 965	Gln	AC)	TC1	CTT Leu	CAC His	Ile	CCA Pro	CCA Pro	CAA Glr	GC1 Ala 975	rras	CTC Leu	Ser	3164
CCT	GAA Glu 980	Ala	TCI Ser	GA?	CTI Lev	TATI Ile 985	· Ile	Lys	CTI Leu	TGC Cys	CG# Arg 990	i er	A CCC	GAA Glu	GAT ABP	3212
Arg 995	Lev	GG(	Lys	AA!	r GG: n Gl: 100	y Ala	GAT A ABI	GAI	ATA 1 11e	100	3 Ale	r CA'	r cci	A TTT	TTT Phe 1010	3260
AA) Lyi	A ACI	A AT	CAC B As	Ph 10	e Se	C AG1 r Sei	C As	CTO	AGI Arg 103	3 GTI	G CAC	G TC n Se	T GC	T TC	A TAC r Tyr 25	3308
ATT	r cc	r AA	A ATO	e Th	A CA r Hi	C CC	A AC	A GAT	p_Th	A TC	A AA' r As	T TT n Ph	T GA e As 10	b sr	r GTT o Val	3356
GA! As	T CC	o As	T AA p Ly 45	A TT s Le	A TG	G AG p Se	r As	T GA P As 50	T AA	C GA	G GA u Gl	u GI	A AA u Aa 55	T GT: n Va	A AAT 1 Asn	3404
GA As	p Th	T CT r Le 60	C AA u As	T GG n Gl	A TG	р ту	T AA r Ly 65	A AA s As	T GG n Gl	A AA y Ly	8 11	T CC .s Pr .70	T GA	A CA u Hi	T GCA s Ala	3452
Ph	C TA e Ty 75	T GA r Gl	A TT u Ph	T AC	ir Ph	c cc le Ar 080	A AG	G TT	T TT e Ph	e as	T GA P As 185	C AF	AT GG	C TA	C CCA r Pro 1090	3500
TA Ty	T AA	T TA	T CC	O L	AG CC 78 P1 095	T AT	T GA e Gl	A TA u Ty	r GI	A TA U Ty .00	c Ai	T AI le Ai	AT TO	27 97	A GGC n Gly	3548



TCA GAG CAG CAG TCG GAT GAA GAT GAT CAA AAC ACA GGC TCA GAG ATT Ser Glu Gln Ser Asp Glu Asp Asp Gln Asn Thr Gly Ser Glu Ile 1110 1115 1120	3596
AAA AAT CGC GAT CTA GTA TAT GTT TAA CACACTAGTA AATAAATGTA Lys Asn Arg Asp Leu Val Tyr Val * 1125 1130	3643
ATGAGGATTT GTAAAAGGGC CTGAAATGCG AGGTGTTTTG AGGTTCTGAG AGTAAAATTA	3703
TGCAAATATG ACAGAGCTAT ATATGTGTGC TCTGTGTACA ATATTTTATT TTCCTAAATT	3763
ATGGGAAATC CTTTTAAAAT GTTAATTTAT TCCAGCCGTT TAAATCAGTA TTTAGAAAAA	3823
AATTGTTATA AGGAAAGTAA ATTATGAACT GAATATTATA GTCAGTTCTT GGTACTTAAA	3883
GTACTTAAAA TAAGTAGTGC TTTGTTTAAA AGGAGAAACC TGGTATCTAT TTGTATATAT	3943
GCTAAATAAT TTTAAAATAC AAGAGTTTTT GAAATTTTTT T	3984

## (2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 1131 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Lys Arg Ser Glu Lys Pro Glu Gly Tyr Arg Gln Met Arg Pro Lys 1 5 10 15

Thr Phe Pro Ala Ser Asn Tyr Thr Val Ser Ser Arg Gln Met Leu Gln 20 25 30

Glu Ile Arg Glu Ser Leu Arg Asn Leu Ser Lys Pro Ser Asp Ala Ala 35 40 45

Lys Ala Glu His Asn Met Ser Lys Met Ser Thr Glu Asp Pro Arg Gln
50 60

Val Arg Asn Pro Pro Lys Phe Gly Thr His His Lys Ala Leu Gln Glu 65 70 75 80

Ile Arg Asn Ser Leu Leu Pro Phe Ala Asn Glu Thr Asn Ser Ser Arg 85 90 95

Ser Thr Ser Glu Val Asn Pro Gln Met Leu Gln Asp Leu Gln Ala Ala 100 105 110

Gly Phe Asp Glu Asp Met Val Ile Gln Ala Leu Gln Lys Thr Asn Asn 115 120 125

Arg Ser Ile Glu Ala Ala Ile Glu Phe Ile Ser Lys Met Ser Tyr Gln 130 135 140

Asp Pro Arg Arg Glu Gln Met Ala Ala Ala Ala Ala Arg Pro Ile Asn 145 150 155 160

Ala Ser Met Lys Pro Gly Asn Val Gln Gln Ser Val Asn Arg Lys Gln 165 170 175

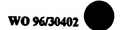
Ser Trp Lys Gly Ser Lys Glu Ser Leu Val Pro Gln Arg His Gly Pro 180 185 190



Pro Leu Gly Glu Ser Val Ala Tyr His Ser Glu Ser Pro Asn Ser Gln 200 205 Thr Asp Val Gly Arg Pro Leu Ser Gly Ser Gly Ile Ser Ala Phe Val Gln Ala His Pro Ser Asn Gly Gln Arg Val Asn Pro Pro Pro Pro Pro Gln Val Arg Ser Val Thr Pro Pro Pro Pro Pro Arg Gly Gln Thr Pro ... 250 255 Pro Pro Arg Gly Thr Thr Pro Pro Pro Pro Ser Trp Glu Pro Asn Ser Gln Thr Lys Arg Tyr Ser Gly Asn Met Glu Tyr Val Ile Ser Arg Ile 280 Ser Pro Val Pro Pro Gly Ala Trp Gln Glu Gly Tyr Pro Pro Pro Pro Leu Asn Thr Ser Pro Het Asn Pro Pro Asn Gln Gly Gln Arg Gly Ile 310 Ser Ser Val Pro Val Gly Arg Gln Pro Ile Ile Met Gln Ser Ser Ser Lys Phe Asn Phe Pro Ser Gly Arg Pro Gly Met Gln Asn Gly Thr Gly Gln Thr Asp Phe Met Ile His Gln Asn Val Val Pro Ala Gly Thr Val Asn Arg Gln Pro Pro Pro Pro Tyr Pro Leu Thr Ala Ala Asn Gly Gln Ser Pro Ser Ala Leu Gln Thr Gly Gly Ser Ala Ala Pro Ser Ser Tyr Thr Asn Gly Ser Ile Pro Gln Ser Met Met Val Pro Asn Arg Asn Ser His Asn Met Glu Leu Tyr Asn Ile Ser Val Pro Gly Leu Gln Thr Asn Trp Pro Gln Ser Ser Ala Pro Ala Gln Ser Ser Pro Ser Ser Gly His Glu Ile Pro Thr Trp Gln Pro Asn Ile Pro Val Arg Ser Asn Ser Phe Asn Asn Pro Leu Gly Asn Arg Ala Ser His Ser Ala Asn Ser Gln Pro Ser Ala Thr Thr Val Thr Ala Ile Thr Pro Ala Pro Ile Gln Gln 490 Pro Val Lys Ser Met Arg Val Leu Lys Pro Glu Leu Gln Thr Ala Leu Ala Pro Thr His Pro Ser Trp Ile Pro Gln Pro Ile Gln Thr Val Gln 515 520 525 Pro Ser Pro Phe Pro Glu Gly Thr Ala Ser Asn Val Thr Val Met Pro Pro Val Ala Glu Ala Pro Asn Tyr Gln Gly Pro Pro Pro Tyr Pro

545					550					555					560
Lys	His	Leu	Leu	His 565	Gln	Asn	Pro	Ser	Val 570	Pro	Pro	Tyr	Glu	Ser 575	Ile
Ser	Lys	Pro	Ser 580	Lys	Glu	Asp	Gln	Pro 585	Ser	Leu	Pro	Lys	Glu 590	Asp	Glu
Ser	Glu	Lys 595	Ser	Tyr	Glu	Asn	Val 600	Asp	Ser	Gly	Asp	Lys 605	Glu	Lys	Lys
Gln	Ile 610	Thr	Thr	Ser	Pro	Ile 615	Thr	Val	Arg	Lys	Asn 620	Lys	Lys	Asp	Glu
Glu 625	Arg	Arg	Glu	Ser	Arg 630	Ile	Gln	Ser	Tyr	Ser 635	Pro	Gln	Ala	Phe	<b>Lув</b> 640
Phe	Phe	Met	Glu	Gln 645	His	Val	Glu	λen	<b>Val</b> 650	Leu	Lys	Ser	His	Gln 655	Gln
Arg	Leu	His	Arg 660	Lys	Гув	Gln	Leu	Glu 665	Asn	Glu	Met	Met	Arg 670	Val	Gly
Leu	Ser	Gln 675	Asp	Ala	Gln	Авр	Gln 680	Met	Arg	Lys	Met	Leu 685	Сув	Gln	Lys
Glu	Ser 690	Asn	Tyr	Ile	Arg	Leu 695	Lys	Arg	Ala	Lув	Met 700	Asp	Lув	Ser	Het
Phe 705	Val	Lys	Ile	Lys	Thr 710	Leu	Gly	Ile	Gly	Ala 715	Phe	Gly	Glu	Val	Сув 720
Leu	Ala	Arg	Lys	Val 725	Asp	Thr	Lye	Ala	<b>Leu</b> 730	Tyr	Ala	Thr	Lys	Thr 735	Leu
Arg	Lys	Lys	740	Val	Leu	Leu	Arg	Авп 745	Gln	Val	Ala	His	Val 750	Lys	Ala
Glu	Arg	<b>Asp</b> 755	Ile	Leu	Ala	Glu	Ala 760	yab	Asn	Glu	Trp	Val 765	Val	Arg	Leu
Tyr	Tyr 770	Ser	Phe	Gln	Asp	<b>Lys</b> 775	Asp	Asn	Leu	Tyr	Phe 780	Val	Met	Asp	Tyr
Ile 785	Pro	Gly	Gly	Asp	Met 790	Met	Ser	Leu	Leu	Ile 795	λrg	Met	Gly	Ile	Phe 800
Pro	Glu	Ser	Leu	Ala 805	Arg	Phe	Tyr	Ile	Ala 810	Glu	Leu	Thr	Сув	Ala 815	Val
Glu	Ser	Val	His 820	Lys	Met	Gly	Phe	11e 825	His	Arg	Asp	Ile	Lys 830	Pro	Asp
Asn	Ile	Leu 835	Ile	Asp	Arg	Asp	Gly 840	His	Ile	Lys		Thr 845	Asp	Phe	Gly
Leu	Сув 850	Thr	Gly	Phe	Arg	Trp 855	Thr	His	Asp	Ser	<b>Lув</b> 860	Tyr	Tyr	Gln	Ser
Gly 865	Asp	His	Pro	Arg	Gln 870	Asp	Ser	Het	Asp	Phe 875	Ser	Asn	Glu	Trp	Gly 880
Asp	Pro	Ser	Ser	Сув 885	Arg	Сув	Gly	Asp	Arg 890	Leu	Lys	Pro		Glu 895	Arg
Arg	Ala	Ala	<b>Arg</b> 900	Gln	His	Gln	Arg	Сув 905	Leu	Ala	His	Ser	<b>Leu</b> 910	Val	Gly

96





Thr Pro Asn Tyr Ile Ala Pro Glu Val Leu Leu Arg Thr Gly Tyr Thr 915 920 925

Gln Leu Cys Asp Trp Trp Ser Val Gly Val Ile Leu Phe Glu Met Leu 930 935 940

Val Gly Gln Pro Pro Phe Leu Ala Gln Thr Pro Leu Glu Thr Gln Met 945 950 955

Lys Val Ile Asn Trp Gln Thr Ser Leu His Ile Pro Pro Gln Ala Lys 965 970 975

Leu Ser Pro Glu Ala Ser Asp Leu Ile Ile Lys Leu Cys Arg Gly Pro 980 985 990

Glu Asp Arg Leu Gly Lys Asn Gly Ala Asp Glu Ile Lys Ala His Pro 995 1000 1005

Phe Phe Lys Thr Ile Asp Phe Ser Ser Asp Leu Arg Gln Gln Ser Ala 1010 1015 1020

Ser Tyr Ile Pro Lys Ile Thr His Pro Thr Asp Thr Ser Asn Phe Asp 1025 1030 1035 1040

Pro Val Asp Pro Asp Lys Leu Trp Ser Asp Asp Asn Glu Glu Asn 1045 1050 1055

Val Asn Asp Thr Leu Asn Gly Trp Tyr Lys Asn Gly Lys His Pro Glu 1060 1065 1070

His Ala Phe Tyr Glu Phe Thr Phe Arg Arg Phe Phe Asp Asp Asn Gly 1075 1080 1085

Tyr Pro Tyr Asn Tyr Pro Lys Pro Ile Glu Tyr Glu Tyr Ile Asn Ser 1090 1095 1100

Gin Gly Ser Glu Gin Gin Ser Asp Glu Asp Asp Gln Asn Thr Gly Ser 1105 1110 1115 1120

Glu Ile Lys Asn Arg Asp Leu Val Tyr Val \* 1125 1130

## (2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 3213 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: cDNA

# (ix) FEATURE:

- (A) NAME/KEY: CDS
- (B) LOCATION: 1..2889

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

GTG CAA CAT TCA ATT AAC CGA AAA CAA AGC TGG AAA GGT TCT AAA GAG
Val Gln His Ser Ile Asn Arg Lys Gln Ser Trp Lys Gly Ser Lys Glu
1 5 10 15

TCT CTA GTT CCT CAG AGA CAC GGC CCA TCT CTA GGA GAA AAT GTG GTT Ser Leu Val Pro Gln Arg His Gly Pro Ser Leu Gly Glu Asn Val 20 25 30

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															CTG Leu	144
															GGA Gly	192
_														ACT Thr		240
														ACT Thr 95		288
														TCT Ser		336
														GCG		384
														ATG Met		432
														GGT Gly		480
CAA Gln	CCC Pro	ATC Ile	ATC Ile	ATG Met 165	CAG Gln	AGT Ser	ACT Thr	AGC Ser	<b>AAA</b> Lys 170	TTT Phe	AAC Asn	TTT Phe	ACA Thr	CCA Pro 175	GGG Gly	528
CGA Arg	CCT Pro	GGA Gly	GTT Val 180	CAG Gln	TAA NBN	GGT Gly	GGT Gly	GGT Gly 185	CAG Gln	TCT Ser	GAT Asp	TTT Phe	ATC Ile 190	GTG Val	CAC Hib	576
														CCA Pro		624
														ACA Thr		672
GCT Ala 225	TCT Ser	GCT Ala	GCT Ala	CCA Pro	CCA Pro 230	TCA Ser	TTC Phe	GCC Ala	AAT Aen	GGA Gly 235	AAC Asn	GTT Val	CCT Pro	CAG Gln	TCG Ser 240	720
ATG Met	ATG Met	GTG Val	CCC Pro	AAC Asn 245	AGG Arg	AAC Asn	AGT Ser	CAT His	AAC Asn 250	ATG Met	GAG Glu	CTT Leu	TAT Tyr	AAT ABN 255	ATT Ile	768
AAT Asn	GTC Val	CCT Pro	GGA Gly 260	CTG Leu	CAA Gln	ACA Thr	GCC Ala	TGG Trp 265	CCC Pro	CAG Gln	TCG Ser	TCT Ser	TCT Ser 270	GCT Ala	CCT Pro	816
GCG Ala	CAG Gln	TCA Ser 275	TCC Ser	CCA Pro	AGC Ser	GGT Gly	GGG Gly 280	CAT His	GAA Glu	ATT Ile	CCT Pro	ACA Thr 285	TGG Trp	CAA Gln	CCT Pro	864
AAC Asn	ATA Ile 290	CCA Pro	GTG Val	AGG Arg	TCA Ser	AAT Asn 295	TCT Ser	TTT Phe	AAT Asn	AAC Asn	CCA Pro 300	TTA Leu	GGA Gly	AGT Ser	AGA Arg	912

GCA Ala 305	AGT Ser	CAC His	TCT Ser	GCT Ala	AAT Asn 310	TCT Ser	CAG Gln	CCT Pro	TCT Ser	GCC Ala 315	ACT Thr	ACA Thr	GTC Val	ACT Thr	GCC Ala 320	960
ATC Ile	ACA Thr	CCC Pro	GCT Ala	CCT Pro 325	ATT Ile	CAA Gln	CAG Gln	CCC Pro	GTG Val 330	AAA Lys	AGC Ser	ATG Met	CGC Arg	GTC Val 335	CTG Leu	1008
AAA Lys	CCA Pro	GAG Glu	CTG Leu 340	CAG Gln	ACT Thr	GCT Ala	TTA Leu	GCC Ala 345	CCA Pro	ACC Thr	CAT His	CCT Pro	TCT Ser 350	TGG Trp	ATG Het	1056
CCA Pro	CAG Gln	CCA Pro 355	GTT Val	CAG Gln	ACT Thr	GTT Val	CAG Gln 360	CCT Pro	ACC Thr	CCT Pro	TTT Phe	TCT Ser 365	GAG Glu	GGT Gly	ACA Thr	1104
				CCT Pro												1152
				CCG Pro												1200
				TAT Tyr 405												1248
				AAG Lys												1296
				AAA Lys												1344
				AAG Lys												1392
				CAG Gln												1440
				TCT Ser 485											CTA Leu	1488
				ATG Met												1536
				CTT Leu												1584
		Lys													GGA Gly	1632
ATA Ile 545	GGA Gly	GCG Ala	TTT Phe	GGT Gly	GAA Glu 550	GTC Val	TGT Cys	CTA Leu	GCA Ala	AGA Arg 555	AAA Lys	GTC Val	GAT Asp	ACT Thr	AAA Lys 560	1680
				ACA Thr 565												1728



AAT	CAG Gln	GTG Val	GCT Ala 580	CAT His	GTG Val	AAA Lys	GCG Ala	GAG Glu 585	AGG Arg	GAT Asp	ATC Ile	CTA Leu	GCA Ala 590	Glu	GCC Ala	1776
GAC Asp	AAT Asn	GAG Glu 595	TGG Trp	GTG Val	GTC Val	CGC Arg	CTG Leu 600	Tyr	TAC Tyr	TCT Ser	TTC	CAG Gln 605	GAC Asp	AAG Lys	GAC Asp	1824
AAC Asn	TTG Leu 610	TAC Tyr	TTT Phe	GTG Val	ATG Het	Asp	TAC	Ile	CCT Pro	GGG Gly	GGG Gly 620	Хвр	Met	ATG Het	AGC Ser	1872
CTA Leu 625	Leu	ATT	AGA Arg	ATG Met	GGC Gly 630	ATC	TTT Phe	CCT	GAA Glu	AAT Asn 635	CTG	GCA Ala	CGA	TTC	TAC Tyr 640	1920
ATA Ile	GCA Ala	GAA Glu	CTT	ACC Thr 645	TGT Cys	GCA Ala	GTT Val	GAA Glu	AGT Ser 650	GTT Val	CAT	AAA Lys	ATG Met	GGT Gly 655	TTT Phe	1968
Ile	His	Arg	GAT Asp 660	Ile	Lys	Pro	Asp	Asn 665	Ile	Leu	Ile	Авр	<b>Arg</b> 670	Asp	Gly	2016
CAT His	ATT	AAA Lys 675	TTG Leu	ACT	GAC Asp	TTT Phe	GGC Gly 680	TTG Leu	TGC Cys	ACT Thr	GGC Gly	TTC Phe 685	AGA Arg	TGG Trp	ACA Thr	2064
CAT His	GAC Asp 690	TCC Ser	AAG Lys	TAC Tyr	TAC Tyr	CAG Gln 695	AGT Ser	GG Gly	GAT Asp	CAC His	CCA Pro 700	CGG Arg	CAA Gln	GAT Asp	AGC Ser	2112
ATG Met 705	GAT Asp	TTC Phe	AGT Ser	AAC Asn	GAA Glu 710	TGG Trp	GGA Gly	GAT Asp	CCT Pro	TCC Ser 715	AAT Asn	TGT Cys	CGG Arg	TGT Cys	GGG Gly 720	2160
Asp	AGA Arg	CTG Leu	AAG Lys	CCA Pro 725	CTG Leu	GAG Glu	CGG Arg	AGA Arg	GCT Ala 730	GCT Ala	CGC Arg	CAG Gln	CAC His	CAG Gln 735	CGA Arg	2208
TGT Cys	CTA Leu	GCC Ala	CAT His 740	TCT Ser	CTG Leu	GTT Val	GGG Gly	ACT Thr 745	CCC Pro	AAT Asn	TAT Tyr	ATT	GCA Ala 750	CCT Pro	GAA Glu	2256
GTG Val	CTA Leu	CTG Leu 755	CGA Arg	ACA Thr	GGA Gly	TAT Tyr	ACA Thr 760	CAG Gln	CTG Leu	TGT Cys	GAC Asp	TGG Trp 765	TGG Trp	AGT Ser	GTT Val	2304
GGT Gly	GTT Val 770	ATT Ile	CTT Leu	TGT Cys	GAA Glu	ATG Met 775	TTG Leu	GTG Val	GGA Gly	CAA Gln	CCT Pro 780	CCT Pro	TTC Phe	TTG Leu	GCA Ala	2352
CAA Gln 785	ACC Thr	CCA Pro	TTA Leu	GAA Glu	ACA Thr 790	CAA Gln	ATG Met	AAG Lys	GTT Val	ATC Ile 795	ATC Ile	TGG Trp	CAA Gln	ACT Thr	TCT Ser 800	2400
CTA Leu	CAC His	ATC Ile	CCT Pro	CCT Pro 805	CAA Gln	GCT Ala	AAG Lyb	CTG Leu	AGT Ser 810	CCT Pro	GAA Glu	GCC Ala	TCT Ser	GAC Asp 815	CTC Leu	2448
ATT Ile	ATC Ile	AAA Lys	CTG Leu 820	TGT Cyb	CGA Arg	GGA Gly	CCA Pro	GAA Glu 825	GAC Asp	CGC Arg	CTC Leu	GGC Gly	AAG Lys 830	AAC Asn	GGT Gly	2496
GCT Ala	GAT Asp	GAG Glu 835	ATA Ile	AAG Lys	GCT Ala	His	CCA Pro 840	TTT Phe	TTT Phe	AAG Lys	ACC Thr	ATC Ile 845	GAT Asp	TTC Phe	TCT Ser	2544



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AGT GAT CTG AGA CAG CAG TCT GCT TCA TAC ATC CCT AAA ATC ACG CAT Ser Asp Leu Arg Gln Gln Ser Ala Ser Tyr Ile Pro Lys Ile Thr His 850 855 860	2592
CCA ACA GAT ACA TCC AAT TTC GAC CCT GTT GAT CCT GAT AAA TTG TGG Pro Thr Asp Thr Ser Asn Phe Asp Pro Val Asp Pro Asp Lys Leu Trp 865 870 875 880	2640
AGC GAT GGC AGC GAG GAA AAT ATC AGT GAC ACT CTG AGC GGA TGG Ser Asp Gly Ser Glu Glu Asn Ile Ser Asp Thr Leu Ser Gly Trp 885	2688
TAT AAA AAT GGG AAG CAC CCC GAG CAC GCT TTC TAT GAG TTC ACC TTT Tyr Lys Asn Gly Lys His Pro Glu His Ala Phe Tyr Glu Phe Thr Phe 900 905	2736
CGG AGG TTT TTT GAT GAC AAT GGC TAC CCA TAT AAT TAT CCA AAG CCT Arg Arg Phe Phe Asp Asp Asn Gly Tyr Pro Tyr Asn Tyr Pro Lys Pro 915	2784
ATT GAG TAT GAA TAC ATT CAT TCA CAG GGC TCA GAA CAA CAG TCT GAT Ile Glu Tyr Glu Tyr Ile His Ser Gln Gly Ser Glu Gln Gln Ser Asp 930 940	2832
GAA GAT GAT CAA CAC ACA AGC TCC GAT GGA AAC AAC CGA GAT CTA GTG Glu Asp Asp Gln His Thr Ser Ser Asp Gly Asn Asn Arg Asp Leu Val 945 955 960	2880
TAT GTT TAA TAAACTAGGA GATCATTGTA AGAATTTGCA AGAGGCCTGA Tyr Val *	2929
AGTGCAGGGG TTTTTGAGAT TTTGAGATAA TTATGCATAT GTGACAGAGT TTGTGTGCTC	2989
TGTGTACAAT ATTTTATTTT CCTAAGTTAT GGGAAATTGT TTTAAAATGT TAATTTATTC	3049
CACCCTTTTA ATTCAGTAAT TTAGAAAAAA TTGTTATAAG GAAAGTAAAT TATGAACTGA	3109
GTATTATAGT CAATTCTTGG TACTTAAAGT ACTTAAAAAG AGAAGCCTGG TATCTTTTGT	3169
ATATATAA AATAATTTTA AAATCCCAAA AAAAAAAAA	3213

## (2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 963 amino acids

  - (B) TYPE: amino acid (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Val Gln His Ser Ile Asn Arg Lys Gln Ser Trp Lys Gly Ser Lys Glu

Ser Leu Val Pro Gln Arg His Gly Pro Ser Leu Gly Glu Asn Val Val 20 25 30

Tyr Arg Ser Glu Ser Pro Asn Ser Gln Ala Asp Val Gly Arg Pro Leu

Ser Gly Ser Gly Ile Ala Ala Phe Ala Gln Ala His Pro Ser Asn Gly 50 60

Gln Arg Val Asn Pro Pro Pro Pro Pro Gln Val Arg Ser Val Thr Pro

65 70 75 Pro Pro Pro Pro Arg Gly Gln Thr Pro Pro Pro Arg Gly Thr Thr Pro Pro Pro Pro Ser Trp Glu Pro Ser Ser Gln Thr Lys Arg Tyr Ser Gly Asn Met Glu Tyr Val Ile Ser Arg Ile Ser Pro Val Pro Pro Gly Ala Trp Gln Glu Gly Tyr Pro Pro Pro Pro Leu Thr Thr Ser Pro Met Asn Pro Pro Ser Gln Ala Gln Arg Ala Ile Ser Ser Val Pro Val Gly Arg Gln Pro Ile Ile Met Gln Ser Thr Ser Lys Phe Asn Phe Thr Pro Gly Arg Pro Gly Val Gln Asn Gly Gly Gly Gln Ser Asp Phe Ile Val His Gln Asn Val Pro Thr Gly Ser Val Thr Arg Gln Pro Pro Pro Tyr 200 Pro Leu Thr Pro Ala Asn Gly Gln Ser Pro Ser Ala Leu Gln Thr Gly Ala Ser Ala Ala Pro Pro Ser Phe Ala Asn Gly Asn Val Pro Gln Ser Met Het Val Pro Asn Arg Asn Ser His Asn Het Glu Leu Tyr Asn Ile Asn Val Pro Gly Leu Gln Thr Ala Trp Pro Gln Ser Ser Ala Pro Ala Gln Ser Ser Pro Ser Gly Gly His Glu Ile Pro Thr Trp Gln Pro Asn Ile Pro Val Arg Ser Asn Ser Phe Asn Asn Pro Leu Gly Ser Arg 295 300 Ala Ser His Ser Ala Asn Ser Gln Pro Ser Ala Thr Thr Val Thr Ala Ile Thr Pro Ala Pro Ile Gln Gln Pro Val Lys Ser Met Arg Val Leu 330 Lys Pro Glu Leu Gln Thr Ala Leu Ala Pro Thr His Pro Ser Trp Met

Pro Gln Pro Val Gln Thr Val Gln Pro Thr Pro Phe Ser Glu Gly Thr 365

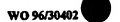
Ala Ser Ser Val Pro Val Ile Pro Pro Val Ala Glu Ala Pro Ser Tyr 370

Gln Gly Pro Pro Pro Pro Pro Pro Lys His Leu Leu His Gln Asn Pro 385

345

Ser Val Pro Pro Tyr Glu Ser Val Ser Lys Pro Cys Lys Asp Glu Gln 405 410 415

Pro Ser Leu Pro Lys Glu Asp Asp Ser Glu Lys Ser Ala Asp Ser Gly
420
425



Asp Ser Gly Asp Lys Glu Lys Lys Gln Ile Thr Thr Ser Pro Ile Thr 440 Val Arg Lys Asn Lys Lys Asp Glu Glu Arg Arg Glu Ser Arg Ile Gln Ser Tyr Ser Pro Gln Ala Phe Lys Phe Phe Met Glu Gln His Val Glu Asn Val Leu Lys Ser His Gln Gln Arg Leu His Arg Lys Lys Gln Leu 490 Glu Asn Glu Met Met Arg Val Gly Leu Ser Gln Asp Ala Gln Asp Gln Het Arg Lys Het Leu Cys Gln Lys Glu Ser Asn Tyr Ile Arg Leu Lys Arg Ala Lys Met Asp Lys Ser Met Phe Val Lys Ile Lys Thr Leu Gly Ile Gly Ala Phe Gly Glu Val Cys Leu Ala Arg Lys Val Asp Thr Lys Ala Leu Tyr Ala Thr Lys Thr Leu Arg Lys Lys Asp Val Leu Leu Arg Asn Gln Val Ala His Val Lys Ala Glu Arg Asp Ile Leu Ala Glu Ala Asp Asn Glu Trp Val Val Arg Leu Tyr Tyr Ser Phe Gln Asp Lys Asp 600 Asn Leu Tyr Phe Val Met Asp Tyr Ile Pro Gly Gly Asp Met Met Ser Leu Leu Ile Arg Met Gly Ile Phe Pro Glu Asn Leu Ala Arg Phe Tyr Ile Ala Glu Leu Thr Cys Ala Val Glu Ser Val His Lys Met Gly Phe 650 Ile His Arg Asp Ile Lys Pro Asp Asn Ile Leu Ile Asp Arg Asp Gly 665 His Ile Lys Leu Thr Asp Phe Gly Leu Cys Thr Gly Phe Arg Trp Thr 675 680 685 His Asp Ser Lys Tyr Tyr Gln Ser Gly Asp His Pro Arg Gln Asp Ser 700 Met Asp Phe Ser Asn Glu Trp Gly Asp Pro Ser Asn Cys Arg Cys Gly Asp Arg Leu Lys Pro Leu Glu Arg Arg Ala Ala Arg Gln His Gln Arg Cys Leu Ala His Ser Leu Val Gly Thr Pro Asn Tyr Ile Ala Pro Glu Val Leu Leu Arg Thr Gly Tyr Thr Gln Leu Cys Asp Trp Trp Ser Val Gly Val Ile Leu Cys Glu Met Leu Val Gly Gln Pro Pro Phe Leu Ala

Gln Thr Pro Leu Glu Thr Gln Met Lys Val Ile Ile Trp Gln Thr Ser



785 790 795 800 Leu His Ile Pro Pro Gln Ala Lys Leu Ser Pro Glu Ala Ser Asp Leu Ile Ile Lys Leu Cys Arg Gly Pro Glu Asp Arg Leu Gly Lys Asn Gly 825 Ala Asp Glu Ile Lys Ala His Pro Phe Phe Lys Thr Ile Asp Phe Ser Ser Asp Leu Arg Gln Gln Ser Ala Ser Tyr Ile Pro Lys Ile Thr His 855 Pro Thr Asp Thr Ser Asn Phe Asp Pro Val Asp Pro Asp Lys Leu Trp 865 870 Ser Asp Gly Ser Glu Glu Asn Ile Ser Asp Thr Leu Ser Gly Trp Tyr Lys Asn Gly Lys His Pro Glu His Ala Phe Tyr Glu Phe Thr Phe Arg Arg Phe Phe Asp Asp Asn Gly Tyr Pro Tyr Asn Tyr Pro Lys Pro 920 925 Ile Glu Tyr Glu Tyr Ile His Ser Gln Gly Ser Glu Gln Gln Ser Asp 930 Glu Asp Asp Gln His Thr Ser Ser Asp Gly Asn Asn Arg Asp Leu Val 950 955 Tyr Val \*

## (2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 3155 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 1..2943

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

ATG Met 1	AGA Arg	GCC Ala	ACC Thr	CCG Pro 5	AAG Lys	TTT	GGA Gly	CCT Pro	TAT Tyr 10	CAA Gln	AAA Lys	GCT Ala	CTC Leu	AGG Arg 15	GAA Glu	48
ATC Ile	CGA Arg	TAT Tyr	TCC Ser 20	CTC Leu	CTG Leu	CCT Pro	TTT Phe	GCC Ala 25	AAC Asn	GAG Glu	TCA Ser	GGC Gly	ACT Thr 30	TCG Ser	GCA Ala	96
GCT Ala	GCA Ala	GAG Glu 35	GTG Val	AAC Asn	CGG Arg	CAG Gln	ATG Met 40	CTT Leu	CAG Gln	GAG Glu	TTG Leu	GTG Val 45	AAT Asn	GCG Ala	GCA Ala	144
TGT Cys	GAC Asp	CAG Gln	GAG Glu	ATG Met	GCT Ala	GGC Gly	AGA Arg	GCG Ala	CTC Leu	ACG Thr	CAG Gln	ACG Thr	GGC Gly	AGT Ser	AGG Arg	192

	50					55					60					
AGT Ser 65	ATC Ile	GAA Glu	GCT Ala	GCC Ala	TTG Leu 70	GAG Glu	TAC Tyr	ATC Ile	AGT Ser	AAG Lys 75	ATG Met	GGC Gly	TAC Tyr	CTG Leu	GAC Asp 80	240
CCC Pro	AGG Arg	AAT Asn	GAG Glu	CAG Gln 85	ATT Ile	GTG Val	CGA Arg	GTC Val	ATC Ile 90	AAG Lys	CAG Gln	ACC Thr	TCC Ser	CCA Pro 95	GGA Gly	288
								ACT Thr 105					-			336
								CAC His								384
								ATG Met								432
								ACC Thr								480
								GCA Ala								528
								CAT His 185								576
								TTC Phe								624
								CAG Gln								672
								TAC Tyr								720
								CCA Pro								768
				Thr				TCT Ser 265							CCG Pro	816
			Ser					TTG Leu								864
		Ser						CGC							_	912
								GTG Val								960
AGG	ACC	AAC	TCC	TTC	AAC	AAC	CCA	CAA	CCT	GAG	ccc	TCA	CTG	ccc	GCC	1008



Arg	Thr	Asn	Ser	Phe 325	Asn	Asn	Pro	Gln	Pro 330		Pro	Ser	Leu	Pro 335	Ala	
Pro	AAC Asn	ACG Thr	GTC Val 340	Thr	GCC Ala	GTG Val	ACG Thr	GCC Ala 345	Ala	CAC	ATC	CTT Leu	CAC His 350	Pro	GTG Val	1056
AAG Lys	AGC Ser	GTG Val 355	CGT	GTG Val	CTG Leu	CGG Arg	CCC Pro 360	GAG Glu	CCC Pro	CAG Gln	ACA Thr	GCC Ala 365	Val	GGG Gly	CCC Pro	1104
TCG Ser	CAC His 370	Pro	GCC Ala	TGG Trp	GTG Val	GCT Ala 375	GCG Ala	CCC	ACA Thr	GCA Ala	CCT Pro 380	GCC Ala	ACT Thr	GAG Glu	AGC Ser	1152
CTG Leu 385	Glu	ACG Thr	AAG Lys	GAG Glu	GGC Gly 390	AGC Ser	GCA Ala	GGC Gly	CCA Pro	CAC His 395	CCG Pro	CTG Leu	Asp Asp	GTG Val	GAC Asp 400	1200
TAT Tyr	GCC	GGC	TCC Ser	GAG Glu 405	CGC Arg	AGG Arg	TGC Cys	CCA Pro	CCG Pro 410	CCT Pro	CCG Pro	TAT Tyr	CCA Pro	AAG Lys 415	CAC His	1248
TTG Leu	CTG Leu	CTG Leu	CCC Pro 420	AGT Ser	AAG Lys	TCT	GAG Glu	CAG Gln 425	TAC Tyr	AGC Ser	GTG Val	GAC Asp	CTG Leu 430	GAC Asp	AGC Ser	1296
CTG Leu	TGC Cys	ACC Thr 435	AGT Ser	GTG Val	CAG Gln	CAG Gln	AGT Ser 440	CTG Leu	CGA Arg	GGG Gly	GGC Gly	ACT Thr 445	GAT Asp	CTA Leu	GAC Asp	1344
GCG	AGT Ser 450	GAC Asp	AAG Lys	AGC Ser	CAC His	AAA Lys 455	GGT Gly	GCG Ala	AAG Lys	GGA Gly	GAC Asp 460	AAA Lys	GCT Ala	GGC	AGA Arg	1392
GAC Asp 465	AAA Lys	AAG Lys	CAG Gln	ATT Ile	CAG Gln 470	ACC Thr	TCC Ser	CCG Pro	GTG Val	CCT Pro 475	GTC Val	CGC Arg	AAG Lys	AAT ABN	AGC Ser 480	1440
AGA Arg	GAT Asp	GAA Glu	GAG Glu	AAG Lys 485	AGA Arg	GAG Glu	TCT Ser	CGC Arg	ATC Ile 490	AAG Lys	AGT Ser	TAC Tyr	TCC Ser	CCT Pro 495	TAT Tyr	1488
GCC Ala	TTC Phe	AAA Lys	TTC Phe 500	TTC Phe	ATG Met	GAG Glu	CAA Gln	CAC His 505	GTG Val	GAG Glu	AAT Asn	GTC Val	ATC Ile 510	AAA Lys	ACC Thr	. 1536
TAC Tyr	CAG Gln	CAG Gln 515	AAG Lys	GTC Val	AGC Ser	CGG Arg	AGG Arg 520	CTA Leu	CAG Gln	CTG Leu	GAG Glu	CAG Gln 525	GAA Glu	ATG Met	GCC Ala	1584
AAA Lys	GCT Ala 530	GGG Gly	CTC Leu	TGT Cys	GAG Glu	GCC Ala 535	GAG Glu	CAG Gln	GAG Glu	CAG Gln	ATG Met 540	AGG Arg	AAG Lys	ATC Ile	CTC Leu	1632
TAC Tyr 545	CAG Gln	AAG Lys	GAG Glu	TCT Ser	AAC Asn 550	TAC Tyr	AAC Asn	CGG Arg	CTG Leu	AAG Lyb 555	AGG Arg	GCC Ala	AAG Lyb	ATG Met	GAC Asp 560	1680
AAG Lys	TCC Ser	ATG Met	Phe	GTG Val 565	AAA Lys	ATC . Ile	AAG Lys	ACT Thr	CTA Leu 570	GGC Gly	ATC Ile	GGT Gly	GCC Ala	TTT Phe 575	GGG Gly	1728
GAA Glu	GTG Val	Сув	CTC Leu 580	GCT Ala	TGT Cys	AAG Lys	Leu	GAC Asp 585	ACT Thr	CAC Hib	GCT Ala	CTG Leu	TAC Tyr 590	GCC Ala	ATG Met	1776

AAG	) )	CTC	<b>N</b> GG	AAG	AAG	GAT	GTC	CTG	AAC	CGG .	AAT	CAA	GTG	GCC	CAT		1824
Lys	Thr	<b>Leu</b> 595	Arg	Lys	Lys	Asp	Val 600	Leu	Asn	Arg	yeu	Gln 605	Val	Ala	His		
GTC Val	AAG Lys 610	GCT Ala	GAG Glu	AGG Arg	GAC Asp	ATC Ile 615	CTG L <b>e</b> u	GCT Ala	GAA Glu	GCA Ala	GAC Asp 620	TAA ABn	GAG Glu	TGG Trp	GTG Val		1872
Val	AAA Lys	Leu	TAC Tyr	TAC Tyr	TCC Ser 630	TTC Phe	CAG Gln	GAC Asp	AAG Lys	GAC Asp 635	AGC Ser	CTG Leu	TAC Tyr	TTT Phe	GTG Val 640		1920
A TOC	GAC	ጥልሮ	ATA Ile	CCA Pro 645	GGC Gly	GGG Gly	GAT Asp	ATG	ATG Het 650	AGC Ser	CTG Leu	CTG Leu	ATC Ile	AGG Arg 655	ATG Het		1968
GAG Glu	GTC Val	TTC Phe	CCT Pro 660	GAG Glu	CAC His	CTG Leu	GCC Ala	CGC Arg 665	TTC Phe	TAC Tyr	ATT Ile	GCA Ala	GAG Glu 670	TTG Leu	ACC Thr		2016
CTG Leu	GCC Ala	ATT Ile 675	GAA Glu	AGT Ser	GTC Val	CAC His	AAG Lys 680	ATG Met	GC	TTT Phe	ATC Ile	CAC His 685	CGG Arg	GAC Asp	ATC Ile		2064
AAG Lys	CCT Pro 690	Asp	AAC Asn	ATA Ile	CTC Leu	ATC Ile 695	GAC Asp	CTG Leu	GAT Asp	GGT Gly	CAT His 700	He	AAG Lys	CTG Leu	ACA Thr		2112
GAT Asp 705	Phe	GGC Gly	CTC Leu	TGC Cys	ACT Thr 710	Gly	TTC Phe	AGG Arg	TGG Trp	ACT Thr 715	CAC His	TAA neA	TCC Ser	AAG Lys	TAC Tyr 720		2160
TAC Tyr	CAG Gln	Lys	GGG	AAC Asn 725	His	ATG Met	AGA Arg	CAG Gln	GAC Asp 730	Ser	ATG Met	GAG Glu	CCC Pro	GGT Gly 735	GAC Asp		2208
CTC	TGG Trp	GAC Asp	GAT Asp 740	Val	TCC	AAC Asn	TGT Cys	CGC Arg 745	Cys	GGA Gly	GAC	AGG Arg	TTA Leu 750	LAB	ACC		2256
CTC	GAG Glu	Glr 755	Arg	GCG Ala	CAG Glr	AAG Lys	Glr 760	His	CAG Glr	AGG Arg	TGC	CTG Leu 765	Ala	CAT His	TCT		2304
CT	GTC Val	Gly	ACI Thi	CCF	AA1	TAC 1 Tyr 775	: Ile	GCT Ala	CCC Pro	GAG Glu	GTC Val 780	r rec	CTC Lev	CGC Arg	Lys		2352
GG( G1; 78	y Ty	C ACC	G CAC	G CTO	TG1 1 Cy1 790	B ASI	TG(	TGG Tr	S AGO	GTC Val 795	GI	r GTC y Val	ATI	CTC Lev	Phe 800		2400
GA: Gl:	G ATG	G CT	G GT u Va	r GG(	y Gl	G CCC	CC	TTO Pho	C TTC E Lei 810	n WIS	C CCG	C ACC	C CCC	Thi 81	A GAG Glu		2448
AC Th	G CA	G CT n Le	G AA u Ly 82	s Va	G ATO	C AA( e As:	C TG	G GA P G1: 82:	u Se	C ACC	G CTO	G CA' u Hi	B 116	e Pri	r ACG o Thr		2496
CA G1	G GT n Va	G AG 1 Ar 83	g Le	C AG u Se	C GC r Al	T GA	G GC u Al 84	a Ar	A GA	C CTO	C AT	C AC e Th 84	r by	G CT s Le	G TGC. u Cyb	•	2544
TG Cy	C GC B Al 85	a Al	T GA a As	C TG p Cy	C CG	C CT g Le 85	u Gl	C AG y Ar	g As	T GG p Gl	G GC y Al 86	a AB	T GA p As	C CT p <b>Le</b>	C AAG u Lyb		2592

## PCT/US96/04101

GCA Ala 865	CAC His	CCG Pro	TTC Phe	TTC Phe	AAC Asn 870	ACC Thr	ATC Ile	GAC Asp	TTT Phe	TCC Ser 875	CGT Arg	GAC Asp	ATC Ile	CGA Arg	AAG Lys 880	2640
	GCT Ala															2688
AAT Asn	TTT Phe	GAC Asp	CCG Pro 900	GTG Val	GAT Asp	GAA Glu	GAA Glu	AGC Ser 905	CCC Pro	TGG Trp	CAC His	GAG Glu	GCC Ala 910	AGC Ser	GGA Gly	2736
GAG Glu	AGC Ser	GCC Ala 915	AAG Lys	GCC Ala	TGG Trp	Asp	ACG Thr 920	CTG Leu	GCC Ala	TCC Ser	CCC Pro	AGC Ser 925	AGC Ser	AAG Lys	CAT His	2784
CCA Pro	GAG Glu 930	CAC His	GCC Ala	TTC Phe	TAT Tyr	GAG Glu 935	TTC Phe	ACC Thr	TTC Phe	CGC Arg	AGG Arg 940	TTC Phe	TTC Phe	GAT Asp	GAC Asp	2832
AAC Asn 945	GGC Gly	TAT Tyr	CCC Pro	TTC Phe	CGG Arg 950	TGC Cys	CCG Pro	AAG Lys	CCC Pro	TCA Ser 955	GAG Glu	CCC Pro	GCA Ala	GAG Glu	AGT Ser 960	2880
GCA Ala	GAC Asp	CCA Pro	GGG Gly	GAT Asp 965	GCG Ala	GAC Asp	TTG Leu	Glu	GGT Gly 970	GCG Ala	GCC Ala	GAG Glu	GGC Gly	TGC Cys 975	CAG Gln	2928
CCG Pro	GTG Val	TAC Tyr	GTG Val 980	TAA *	GCCT	CAGT	TA A	CCAC	AACI	C GA	GGAA	ACCO	AAA	ATGA	GAT	2983
TTCI	TTTC	AG A	AGAC	AAAC	T CA	AGCT	TAGG	AAT	CCTI	CAT	TTTT	AGTT	CT G	GTAA	ATGGG	3043
CAAC	AGGA	AG A	GTCA	ACAT	G AT	TTCA	AATT	AGC	CCTC	TGA	GGAC	CTTC	AC I	GCAT	TAAAA	3103
CAGT	ATTT	TT T	'AAAA'	AATT	A GT	ACAG	TATG	GAA	AGAG	CAC	TTAT	TTTG	GG G	G		3155

# (2) INFORMATION FOR SEQ ID NO:8:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 981 amino acids

  - (B) TYPE: amino acid (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Het Arg Ala Thr Pro Lys Phe Gly Pro Tyr Gln Lys Ala Leu Arg Glu

Ile Arg Tyr Ser Leu Leu Pro Phe Ala Asn Glu Ser Gly Thr Ser Ala

Ala Ala Glu Val Asn Arg Gln Met Leu Gln Glu Leu Val Asn Ala Ala

Cys Asp Gln Glu Met Ala Gly Arg Ala Leu Thr Gln Thr Gly Ser Arg
50 55 60

Ser Ile Glu Ala Ala Leu Glu Tyr Ile Ser Lys Met Gly Tyr Leu Asp 65 70 75 80

Pro Arg Asn Glu Gln Ile Val Arg Val Ile Lys Gln Thr Ser Pro Gly

85 90 95

Lys Gly Leu Ala Ser Thr Pro Val Thr Arg Arg Pro Ser Phe Glu Gly Thr Gly Glu Ala Leu Pro Ser Tyr His Gln Leu Gly Gly Ala Asn Tyr Glu Gly Pro Ala Ala Leu Glu Glu Met Pro Arg Gln Tyr Leu Asp Phe Leu Phe Pro Gly Ala Gly Ala Gly Thr His Gly Ala Gln Ala His Gln 145 150 155 160 His Pro Pro Lys Gly Tyr Ser Thr Ala Val Glu Pro Ser Ala His Phe Pro Gly Thr His Tyr Gly Arg Gly His Leu Leu Ser Glu Gln Ser Gly Tyr Gly Val Gln Arg Ser Ser Ser Phe Gln Asn Lys Thr Pro Pro Asp Ala Tyr Ser Ser Met Ala Lys Ala Gln Gly Gly Pro Pro Ala Ser Leu 215 Thr Phe Pro Ala His Ala Gly Leu Tyr Thr Ala Ser His His Lys Pro Ala Ala Thr Pro Pro Gly Ala His Pro Leu His Val Leu Gly Thr Arg Gly Pro Thr Phe Thr Gly Glu Ser Ser Ala Gln Ala Val Leu Ala Pro Ser Arg Asn Ser Leu Asn Ala Asp Leu Tyr Glu Leu Gly Ser Thr Val Pro Trp Ser Ala Ala Pro Leu Ala Arg Arg Asp Ser Leu Gln Lys Gln 295 Gly Leu Glu Ala Ser Arg Pro His Val Ala Phe Arg Ala Gly Pro Ser Arg Thr Asn Ser Phe Asn Asn Pro Gln Pro Glu Pro Ser Leu Pro Ala 330 Pro Asn Thr Val Thr Ala Val Thr Ala Ala His Ile Leu His Pro Val Lys Ser Val Arg Val Leu Arg Pro Glu Pro Gln Thr Ala Val Gly Pro Ser His Pro Ala Trp Val Ala Ala Pro Thr Ala Pro Ala Thr Glu Ser Leu Glu Thr Lys Glu Gly Ser Ala Gly Pro His Pro Leu Asp Val Asp 395 Tyr Gly Gly Ser Glu Arg Arg Cys Pro Pro Pro Tyr Pro Lys His 410 Leu Leu Pro Ser Lys Ser Glu Gln Tyr Ser Val Asp Leu Asp Ser Leu Cys Thr Ser Val Gln Gln Ser Leu Arg Gly Gly Thr Asp Leu Asp 440



Gly Ser Asp Lys Ser His Lys Gly Ala Lys Gly Asp Lys Ala Gly Arg 455 Asp Lys Lys Gln Ile Gln Thr Ser Pro Val Pro Val Arg Lys Asn Ser Arg Asp Glu Glu Lys Arg Glu Ser Arg Ile Lys Ser Tyr Ser Pro Tyr Ala Phe Lys Phe Phe Met Glu Gln His Val Glu Asn Val Ile Lys Thr 500 505 Tyr Gln Gln Lys Val Ser Arg Arg Leu Gln Leu Glu Gln Glu Het Ala Lys Ala Gly Leu Cys Glu Ala Glu Gln Glu Gln Het Arg Lys Ile Leu Tyr Gln Lys Glu Ser Asn Tyr Asn Arg Leu Lys Arg Ala Lys Met Asp Lys Ser Met Phe Val Lys Ile Lys Thr Leu Gly Ile Gly Ala Phe Gly Glu Val Cys Leu Ala Cys Lys Leu Asp Thr His Ala Leu Tyr Ala Met 585 Lys Thr Leu Arg Lys Lys Asp Val Leu Asn Arg Asn Gln Val Ala His 600 Val Lys Ala Glu Arg Asp Ile Leu Ala Glu Ala Asp Asn Glu Trp Val Val Lys Leu Tyr Tyr Ser Phe Gln Asp Lys Asp Ser Leu Tyr Phe Val Met Asp Tyr Ile Pro Gly Gly Asp Met Met Ser Leu Leu Ile Arg Met 650 Glu Val Phe Pro Glu His Leu Ala Arg Phe Tyr Ile Ala Glu Leu Thr 660 665 Leu Ala Ile Glu Ser Val His Lys Met Gly Phe Ile His Arg Asp Ile Lys Pro Asp Asn Ile Leu Ile Asp Leu Asp Gly His Ile Lys Leu Thr Asp Phe Gly Leu Cys Thr Gly Phe Arg Trp Thr His Asn Ser Lys Tyr Tyr Gln Lys Gly Asn His Met Arg Gln Asp Ser Met Glu Pro Gly Asp Leu Trp Asp Asp Val Ser Asn Cys Arg Cys Gly Asp Arg Leu Lys Thr Leu Glu Gln Arg Ala Gln Lys Gln His Gln Arg Cys Leu Ala His Ser Leu Val Gly Thr Pro Asn Tyr Ile Ala Pro Glu Val Leu Leu Arg Lys Gly Tyr Thr Gln Leu Cys Asp Trp Trp Ser Val Gly Val Ile Leu Phe Glu Met Leu Val Gly Gln Pro Pro Phe Leu Ala Pro Thr Pro Thr Glu

805

810

815

Thr Gln Leu Lys Val Ile Asn Trp Glu Ser Thr Leu His Ile Pro Thr 820 825 830

Gln Val Arg Leu Ser Ala Glu Ala Arg Asp Leu Ile Thr Lys Leu Cys 835 840 845

Cys Ala Ala Asp Cys Arg Leu Gly Arg Asp Gly Ala Asp Asp Leu Lys 850 860

Ala His Pro Phe Phe Asn Thr Ile Asp Phe Ser Arg Asp Ile Arg Lys 865 870 875

Gin Ala Ala Pro Tyr Val Pro Thr Ile Ser His Pro Het Asp Thr Ser 885 890 895

Asn Phe Asp Pro Val Asp Glu Glu Ser Pro Trp His Glu Ala Ser Gly 900 905 910

Glu Ser Ala Lys Ala Trp Asp Thr Leu Ala Ser Pro Ser Ser Lys His 915 920 925

Pro Glu His Ala Phe Tyr Glu Phe Thr Phe Arg Arg Phe Phe Asp Asp 930 940

Asn Gly Tyr Pro Phe Arg Cys Pro Lys Pro Ser Glu Pro Ala Glu Ser 945 950 955 960

Ala Asp Pro Gly Asp Ala Asp Leu Glu Gly Ala Ala Glu Gly Cys Gln 970 975

Pro Val Tyr Val

- (2) INFORMATION FOR SEQ ID NO:9:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 6 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Asp Leu Lys Pro Glu Asn

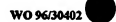
- (2) INFORMATION FOR SEQ ID NO:10:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 9 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:
    - (A) NAME/KEY: Peptide
    - (B) LOCATION: 2
    - (D) OTHER INFORMATION: /label= A

/note= "X at the second position can be either Threonine or



### Serine."

- (ix) FEATURE:
  - (A) NAME/KEY: Peptide
  - (B) LOCATION: 5
- (D) OTHER INFORMATION: /label= B /note= "X at the fifth position can either be Tyrosine or Phenylalanine."
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
  - Gly Xaa Xaa Xaa Xaa Ala Pro Glu 1 5
- (2) INFORMATION FOR SEQ ID NO:11:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 620 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:
  - Met Asp Asn Thr Asn Arg Pro His Leu Asn Leu Gly Thr Asn Asp Thr 1 5 10 15
  - Arg Met Ala Pro Asn Asp Arg Thr Tyr Pro Thr Thr Pro Ser Thr Phe 20 25 30
  - Pro Gln Pro Val Phe Pro Gly Gln Gln Ala Gly Gly Ser Gln Gln Tyr 35 40 45
  - Asn Gln Ala Tyr Ala Gln Ser Gly Asn Tyr Tyr Gln Gln Asn His Asn 50 55 60
  - Asp Pro Asn Thr Gly Leu Ala His Gln Phe Ala His Gln Asn Ile Gly 65 70 75 80
  - Ser Ala Gly Arg Ala Ser Pro Tyr Gly Ser Arg Gly Pro Ser Pro Ala 85 90 95
  - Gln Arg Pro Arg Thr Ser Gly Asn Ser Gly Gln Gln Gln Thr Tyr Gly
    100 105 110
  - Asn Tyr Leu Ser Ala Pro Met Pro Ser Asn Thr Gln Thr Glu Phe Ala 115 120 125
  - Pro Leu Pro Ser Gly Thr Pro Thr Asn Met Ala Pro Met Pro Thr Thr 130 140
  - Thr Arg Arg Ser Ala His Ser Trp Pro Leu Thr Ser Leu Arg Thr Ala 145 150 155 160
  - Ser Ser Ala Pro Gly Ser Ala Thr Arg Gly Glu Cys Cys Ser Asp Ala 165 170 175
  - Leu Leu Pro Leu His Pro Ala Val Ile Gly Ala Asp Thr Leu Phe Arg 180 185 190
  - Gln Ser Glu Met Glu Gln Lys Leu Gly Glu Thr Asn Asp Ala Arg Arg 195 200 205



Arg Glu Ser Ile Trp Ser Thr Ala Gly Arg Lys Glu Gly Gln Tyr Leu 215 Arg Phe Leu Arg Thr Lys Asp Lys Pro Glu Asn Tyr Gln Thr Ile Lys Ile Ile Gly Lys Gly Ala Phe Gly Glu Val Lys Leu Val Gln Lys Lys 245 250 255 Ala Asp Gly Lys Val Tyr Ala Met Lys Ser Leu Ile Lys Thr Glu Met 260 265 270 Phe Lys Lys Asp Gln Leu Ala His Val Arg Ala Glu Arg Asp Ile Leu Ala Glu Ser Asp Ser Pro Trp Val Val Lys Leu Tyr Thr Thr Phe Gln Asp Ala Asn Phe Leu Tyr Met Leu Het Glu Phe Leu Pro Gly Gly Asp 315 Leu Met Thr Met Leu Ile Lys Tyr Glu Ile Phe Ser Glu Asp Ile Thr Arg Phe Tyr Ile Ala Glu Ile Val Leu Ala Ile Asp Ala Val His Lys Leu Gly Phe Ile His Arg Asp Ile Lys Pro Asp Asn Ile Leu Leu Asp Arg Gly Gly His Val Lys Leu Thr Asp Phe Gly Leu Ser Thr Gly Phe His Lys Leu His Asp Asn Asn Tyr Tyr Thr Gln Leu Ceu Gln Gly Lys 385 390 395 395 Ser Asn Lys Pro Arg Asp Asn Arg Asn Ser Val Ala Ile Asp Gin Ile Asn Leu Thr Val Ser Asn Arg Ala Gln Ile Asn Asp Trp Arg Arg Ser 425 Arg Arg Leu Met Ala Tyr Ser Thr Val Gly Thr Pro Asp Tyr Ile Ala 440 Pro Glu Ile Phe Thr Gly His Gly Tyr Ser Phe Asp Cys Asp Trp Trp Ser Leu Gly Thr Ile Met Phe Glu Cys Leu Val Gly Trp Pro Pro Phe Cys Ala Glu Asp Ser His Asp Thr Tyr Arg Lys Ile Val Asn Trp Arg His Ser Leu Tyr Phe Pro Asp Asp Ile Thr Leu Gly Val Asp Ala Glu 505 Asn Leu Ile Arg Ser Leu Ile Cys Asn Thr Glu Asn Arg Leu Gly Arg 520 Gly Gly Ala His Glu Ile Lys Ser His Ala Phe Phe Arg Gly Val Glu Phe Asp Ser Leu Arg Arg Ile Arg Ala Pro Phe Glu Pro Arg Leu Thr 555 Ser Ala Ile Asp Thr Thr Tyr Phe Pro Thr Asp Glu Ile Asp Gln Thr

570

- Asp Asn Ala Thr Leu Leu Lys Ala Gln Gln Ala Ala Arg Gly Ala Ala 580 590
- Ala Pro Ala Gln Gln Glu Glu Ser Pro Glu Leu Ser Leu Pro Phe Ile 595 600 605
- Gly Tyr Thr Phe Lys Arg Phe Asp Asn Asn Phe Arg 610 615 620
- (2) INFORMATION FOR SEQ ID NO:12:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 526 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:
  - Met Asp Ser Ala Arg Gly Trp Phe Gln Lys Leu Ser Ser Thr Lys Lys

    1 10 15
  - Asp Pro Met Ala Ser Gly Arg Glu Asp Gly Lys Pro Val Ser Ala Glu 20 25 30
  - Glu Ala Ser Asn Ile Thr Lys Gln Arg Val Ala Ala Ala Lys Gln Tyr 35 40 45
  - Ile Glu Lys His Tyr Arg Glu Gln Met Lys Asn Leu Gln Glu Arg Arg 50 55 60
  - Glu Arg Arg Ile Leu Leu Glu Lys Lys Leu Ala Asp Ala Asp Val Ser 65 70 75 80
  - Glu Glu Asp Gln Asn Asn Leu Leu Lys Phe Leu Glu Lys Lys Glu Thr 85 90 95
  - Glu Tyr Met Arg Leu Gln Arg His Lys Met Gly Ala Asp Asp Phe Glu 100 105 110
  - Leu Leu Thr Met Ile Gly Lys Gly Ala Phe Gly Glu Pro Ile Cys Met 115 120 125
  - Ile Gly Phe Ser Val Ile Thr Gly Gln Asn Cys Arg Glu Lys Thr Thr 130 140
  - Gly Gln Val Tyr Ala Met Lys Lys Leu Lys Lys Ser Glu Met Leu Arg 145 150 155 160
  - Arg Gly Gln Val Glu His Val Lys Ala Glu Arg Asn Leu Leu Ala Glu 165 170 175
  - Val Asp Ser Asp Cys Ile Val Lys Leu Tyr Tyr Ser Phe Gln Asp Asp 180 185 190
  - Asp Tyr Leu Tyr Leu Val Met Glu Tyr Leu Pro Gly Gly Asp Met Met 195 200 205
  - Thr Leu Leu Met Arg Lys Asp Ile Leu Thr Glu Asp Glu Ala Arg Phe 210 215 220



Tyr Val Ala Glu Thr Val Leu Ala Ile Glu Ser Ile His Lys His Asn 235 Tyr Ile His Arg Asp Ile Lys Pro Asp Asn Leu Leu Leu Asp Arg Tyr Gly His Leu Lys Leu Ser Asp Phe Gly Leu Cys Lys Pro Leu Asp Cys Ser Thr Leu Glu Glu Lys Asp Phe Ser Val Gly Asp Asn Ala Asn Gly Gly Ser Arg Ser Asp Ser Pro Pro Ala Pro Lys Arg Thr Gln Glu Glu Gln Leu Glu His Trp Gln Lys Asn Arg Arg Met Leu Ala Tyr Ser Thr Val Gly Thr Pro Asp Tyr Ile Ala Pro Glu Val Leu Leu Lys Lys Gly 330 Tyr Gly Het Glu Cys Asp Trp Trp Ser Leu Gly Ala Ile Het Tyr Glu Met Leu Val Gly Tyr Pro Pro Phe Tyr Ser Asp Asp Pro Met Ser Thr Cys Arg Lys Ile Val Asn Trp Lys Asn His Leu Lys Phe Pro Glu Glu Ala Lys Leu Ser Pro Glu Ala Lys Asp Ile Ile Ser Arg Leu Leu Cys 395 Asn Val Thr Glu Arg Leu Gly Ser Asn Gly Ala Asp Glu Ile Lys Val His Ser Trp Phe Lys Gly Ile Asp Trp Asp Arg Ile Tyr Gln Met Glu 425 Ala Ala Phe Ile Pro Glu Val Asn Asp Glu Leu Asp Thr Gln Asn Phe Glu Lys Phe Glu Glu Ser Glu Ser His Ser Gln Ser Gly Ser Arg Ser Gly Pro Trp Arg Lys Het Leu Ser Ser Lys Asp Ile Asn Phe Val Gly Tyr Thr Tyr Lys Asn Phe Lys Val Val Asn Asp Tyr Gln Val Pro Gly 490 485 Met Val Glu Leu Lys Lys Thr Asn Thr Lys Pro Lys Lys Pro Thr Ile 505 Lys Ser Leu Phe Gly Asp Glu Ser Glu Ala Ser Glu Asp Asn 515

#### (2) INFORMATION FOR SEQ ID NO:13:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 479 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein

WO 96/30

565

570

575

Asp Asn Ala Thr Leu Leu Lys Ala Gln Gln Ala Ala Arg Gly Ala Ala 580 585 590

Ala Pro Ala Gln Glu Glu Ser Pro Glu Leu Ser Leu Pro Phe Ile 595 600 605

Gly Tyr Thr Phe Lys Arg Phe Asp Asn Asn Phe Arg 610 615 620

#### (2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 526 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Het Asp Ser Ala Arg Gly Trp Phe Gln Lys Leu Ser Ser Thr Lys Lys

1 10 15

Amp Pro Met Ala Ser Gly Arg Glu Amp Gly Lym Pro Val Ser Ala Glu 20 25 30

Glu Ala Ser Asn Ile Thr Lys Gln Arg Val Ala Ala Ala Lys Gln Tyr 35 40 45

Ile Glu Lys His Tyr Arg Glu Gln Met Lys Asn Leu Gln Glu Arg Arg 50 55 60

Glu Arg Arg Ile Leu Leu Glu Lys Lys Leu Ala Asp Ala Asp Val Ser 65 70 75

Glu Glu Asp Gln Asn Asn Leu Leu Lys Phe Leu Glu Lys Lys Glu Thr 85 90 95

Glu Tyr Met Arg Leu Gln Arg His Lys Met Gly Ala Asp Asp Phe Glu 100 105 110

Leu Leu Thr Met Ile Gly Lys Gly Ala Phe Gly Glu Pro Ile Cys Met 115 120 125

Ile Gly Phe Ser Val Ile Thr Gly Gln Asn Cys Arg Glu Lys Thr Thr 130 135 140

Gly Gln Val Tyr Ala Met Lys Lys Leu Lys Lys Ser Glu Met Leu Arg 145 150 155 160

Arg Gly Gln Val Glu His Val Lys Ala Glu Arg Asn Leu Leu Ala Glu 165 170 175

Val Asp Ser Asp Cys Ile Val Lys Leu Tyr Tyr Ser Phe Gln Asp Asp 180 185 190

Asp Tyr Leu Tyr Leu Val Met Glu Tyr Leu Pro Gly Gly Asp Met Met 195 200 205

Thr Leu Leu Het Arg Lys Asp Ile Leu Thr Glu Asp Glu Ala Arg Phe 210 215 220

345

350

Phe Ile Pro Glu Val Asn Asp Glu Leu Asp Thr Gln Asn Phe Glu Lys 355 360 365

Phe Glu Glu Ala Asp Asn Ser Ser Gln Ser Thr Ser Lys Ala Gly Pro 370 375

Trp Arg Lys Met Leu Ser Ser Lys Asp Leu Asn Phe Val Gly Tyr Thr 385 390 395 400

Tyr Lys Asn Phe Glu Ile Val Asn Asp Tyr Gln Val Pro Gly Ile Ala 405 410 415

Glu Leu Lys Lys Asp Thr Lys Pro Lys Arg Pro Ser Ile Lys Ser 420 425 430

Leu Phe Glu Asp Glu Ser Ser Asp Ser Ser Glu Ala Ala Thr Ser Gly
435
440

Asp Gln Ser Val Gln Gly Ser Phe Leu Asn Leu Leu Pro Pro Gln Leu 450 455 460

Glu Val Ser Gln Thr Gln Thr Glu Val Pro Pro Pro Lys Phe Thr 465 470 475

### (2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 500 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Met Glu Lys Val Lys Ala Ala Lys Lys Phe Ile Glu Asn His Tyr Arg 1 5 10 15

Ser Gln Met Lys Asn Ile Gln Glu Arg Lys Glu Arg Arg Trp Val Leu 20 25 30

Glu Lys Gln Leu Ala Ser Ser Asp Val Pro Glu Glu Glu Gln Met Ser 35 40 45

Leu Ile Lys Asp Leu Glu Arg Lys Glu Thr Glu Phe Met Arg Leu Lys 50 60

Arg Asn Arg Ile Cys Val Asn Asp Phe Glu Leu Leu Thr Ile Ile Gly 65 70 80

Arg Gly Ala Tyr Gly Glu Val Gln Leu Cys Arg Glu Lys Lys Ser Glu 85 90 95

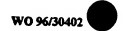
Asn Ile Tyr Ala Met Lys Lys Leu Lys Lys Ser Glu Met Leu Ser Arg 100 105 110

Gly Gln Val Glu His Val Arg Ala Glu Arg Asn Leu Leu Ala Glu Val 115 120 125

Asp Ser His Cys Ile Val Lys Leu Phe Tyr Ser Phe Gln Asp Ala Glu 130 135 140



Tyr Leu Tyr Leu Ile Met Glu Tyr Leu Pro Gly Gly Asp Met Met Thr Leu Leu Met Arg Glu Asp Ile Leu Thr Glu Lys Val Ala Lys Phe Tyr Ile Ala Gln Ser Val Leu Ala Ile Glu Ser Ile His Lys His Asn Tyr Ile His Arg Asp Ile Lys Pro Asp Asn Leu Leu Leu Asp Lys Asn Gly His Met Lys Leu Ser Asp Phe Gly Leu Cys Lys Pro Leu Asp Cys Ala Thr Leu Ser Thr Ile Lys Glu Asn Glu Ser Met Asp Asp Val Ser Lys Asn Ser Het Asp Ile Asp Ala Ser Leu Pro Asp Ala Gly Asn Gly His Ser Trp Arg Ser Ala Arg Glu Gln Leu Gln His Trp Gln Arg Asn Arg Arg Lys Leu Ala Phe Ser Thr Val Gly Thr Pro Asp Tyr Ile Ala Pro Glu Val Leu Leu Lys Lys Gly Tyr Gly Met Glu Cys Asp Trp Trp Ser 290 295 300 Leu Gly Ala Ile Met Tyr Glu Met Leu Val Gly Tyr Pro Pro Phe Tyr Ser Asp Asp Pro Ile Thr Thr Cys Arg Lys Ile Val His Trp Arg His Tyr Leu Lys Phe Pro Asp Asp Ala Lys Leu Thr Phe Glu Ala Arg Asp 340 345 Leu Ile Cys Arg Leu Leu Cys Asp Val Glu His Arg Leu Gly Thr Gly Gly Ala Glu Gln Ile Lys Val His Ala Trp Phe Lys Asp Val Glu Trp Asp Arg Leu Tyr Glu Thr Asp Ala Ala Tyr Lys Pro Gln Val Asn Gly Glu Leu Asp Thr Gln Asn Phe Met Lys Phe Asp Glu Ala Asn Pro Pro Thr Pro Ser Arg Ser Gly Ser Gly Pro Ser Arg Lys Met Leu Thr Ser Lys Asp Leu Ser Phe Val Gly Tyr Thr Tyr Lys Asn Phe Asp Ala Val Lys Gly Leu Lys His Ser Phe Asp Arg Lys Gly Ser Thr Ser Pro Lys Arg Pro Ser Leu Asp Ser Met Phe Asn Glu Asn Gly Met Asp Tyr Thr Ala Lys His Ala Glu Glu Het Asp Val Gln Met Leu Thr Ala Asp Asp 485 490 Cys Met Ser Pro



# (2) INFORMATION FOR SEQ ID NO:15:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 564 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

Met Phe Ser Arg Ser Asp Arg Glu Val Asp Asp Leu Ala Gly Asn Met
1 5 10 15

Ser His Leu Gly Phe Tyr Asp Leu Asn Ile Pro Lys Pro Thr Ser Pro 20 25 30

Gln Ala Gln Tyr Arg Pro Ala Arg Lys Ser Glu Asn Gly Arg Leu Thr 35 40

Pro Gly Leu Pro Arg Ser Tyr Lys Pro Cys Asp Ser Asp Asp Gln Asp 50 55

Thr Phe Lys Asn Arg Ile Ser Leu Asn His Ser Pro Lys Lys Leu Pro 65 70 75 80

Lys Asp Phe His Glu Arg Ala Ser Gln Ser Lys Thr Gln Arg Val Val 85 90 95

Asn Val Cys Gln Leu Tyr Phe Leu Asp Tyr Tyr Cys Asp Met Phe Asp 100 105 110

Tyr Val Ile Ser Arg Arg Gln Arg Thr Lys Gln Val Leu Arg Tyr Leu 115 120 125

Glu Gln Gln Arg Ser Val Lys Asn Val Ser Asn Lys Val Leu Asn Glu 130 135 140

Glu Trp Ala Leu Tyr Leu Gln Arg Glu His Glu Val Leu Arg Lys Arg 145 150 155 160

Arg Leu Lys Pro Lys His Lys Asp Phe Gln Ile Leu Thr Gln Val Gly
165 170 175

Gln Gly Gly Tyr Gly Gln Val Tyr Leu Ala Lys Lys Lys Asp Ser Asp 180 185 190

Glu Ile Cys Ala Leu Lys Ile Leu Asn Lys Lys Leu Leu Phe Lys Leu 195 200 205

Asn Glu Thr Asn His Val Leu Thr Glu Arg Asp Ile Leu Thr Thr 210 215 220

Arg Ser Asp Trp Leu Val Lys Leu Leu Tyr Ala Phe Gln Asp Pro Glu 225 235 240

Ser Leu Tyr Leu Ala Met Glu Phe Val Pro Gly Gly Asp Phe Arg Thr 245 250 255

Leu Leu Ile Asn Thr Arg Ile Leu Lys Ser Gly His Ala Arg Phe Tyr 260 265 270



Ile Ser Glu Het Phe Cys Ala Val Asn Ala Leu His Glu Leu Gly Tyr Thr His Arg Asp Leu Lys Pro Glu Asn Phe Leu Ile Asp Ala Thr Gly His Ile Lys Leu Thr Asp Phe Gly Leu Ala Ala Gly Thr Val Ser Asn Glu Arg Ile Glu Ser Het Lys Ile Arg Leu Glu Glu Val Lys Asn Leu Gln Phe Pro Ala Phe Thr Glu Arg Ser Ile Glu Asp Arg Ser Lys Ile Tyr His Asn Het Arg Lys Thr Glu Ile Asn Tyr Ala Asn Ser Het Val Gly Ser Pro Asp Tyr Het Ala Leu Glu Val Leu Glu Gly Lys Lys Tyr Asp Phe Thr Val Asp Tyr Trp Ser Leu Gly Cys Met Leu Phe Glu Ser 390 395 Leu Val Gly Tyr Thr Pro Phe Ser Gly Ser Ser Thr Asn Glu Thr Tyr Glu Asn Leu Arg Tyr Trp Lys Lys Thr Leu Arg Arg Pro Arg Thr Glu Asp Arg Arg Ala Ala Phe Ser Asp Arg Thr Trp Asp Leu Ile Thr Arg Leu Ile Ala Asp Pro Ile Asn Arg Val Arg Ser Phe Glu Gln Val Arg Lys Met Ser Tyr Phe Ala Glu Ile Asn Phe Glu Thr Leu Arg Thr Ser Ser Pro Pro Phe Ile Pro Gln Leu Asp Asp Glu Thr Asp Ala Gly Tyr Phe Asp Asp Phe Thr Asn Glu Glu Asp Met Ala Lys Tyr Ala Asp Val Phe Lys Arg Gln Asn Lys Leu Ser Ala Met Val Asp Asp Ser Ala Val Asp Ser Lys Leu Val Gly Phe Thr Phe Arg His Arg Asp Gly Lys Gln 535 Gly Ser Ser Gly Ile Leu Tyr Asn Gly Ser Glu His Ser Asp Pro Phe 545

Ser Thr Phe Tyr

# (2) INFORMATION FOR SEQ ID NO:16:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 561 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein



(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Met Ala Gly Asn Met Ser Asn Leu Ser Phe Asp Gly His Gly Thr Pro 1 5 10 15

Gly Gly Thr Gly Leu Phe Pro Asn Gln Asn Ile Thr Lys Arg Arg Thr

Arg Pro Ala Gly Ile Asn Asp Ser Pro Ser Pro Val Lys Pro Ser Phe

Phe Pro Tyr Glu Asp Thr Ser Asn Met Asp Ile Asp Glu Val Ser Gln 50 55 60

Pro Asp Met Asp Val Ser Asn Ser Pro Lys Lys Leu Pro Pro Lys Phe 65 70 75 80

Tyr Glu Arg Ala Thr Ser Asn Lys Thr Gln Arg Val Val Ser Val Cys 85 90 95

Lys Met Tyr Phe Leu Glu Tyr Tyr Cys Asp Met Phe Asp Tyr Val Ile 100 105 110

Ser Arg Arg Gln Arg Thr Lys Gln Val Leu Glu Tyr Leu Gln Gln Gln 115 120 125

Ser Gln Leu Pro Asn Ser Asp Gln Ile Lys Leu Asn Glu Glu Trp Ser 130 135 140

Ser Tyr Leu Gln Arg Glu His Gln Val Leu Arg Lys Arg Arg Leu Lys 150 155 160

Pro Lys Asn Arg Asp Phe Glu Met Ile Thr Gln Val Gly Gln Gly Gly 165 170 175

Tyr Gly Gln Val Tyr Leu Ala Arg Lys Lys Asp Thr Lys Glu Val Cys 180 185 190

Ala Leu Lys Ile Leu Asn Lys Lys Leu Leu Phe Lys Leu Asn Glu Thr 195 200 205

Lys His Val Leu Thr Glu Arg Asp Ile Leu Thr Thr Arg Ser Glu 210 215 220

Trp Leu Val Lys Leu Leu Tyr Ala Phe Gln Glu Leu Gln Ser Leu Tyr 225 230 235 240

Leu Ala Met Glu Phe Val Pro Gly Gly Asp Phe Arg Thr Leu Leu Ile 245 250 255

Asn Thr Arg Cys Leu Lys Ser Gly His Ala Arg Phe Tyr Ile Ser Glu 260 265 270

Met Phe Cys Ala Val Asn Ala Leu His Asp Leu Gly Tyr Thr His Arg 275 280 285

Asp Leu Lys Pro Glu Asn Phe Leu Ile Asp Ala Lys Gly His Ile Lys 290 295 300

Leu Thr Asp Phe Gly Leu Ala Ala Gly Thr Ile Ser Asn Glu Arg Ile 305 310 315

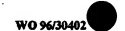
Glu Ser Met Lys Ile Arg Leu Glu Lys Ile Lys Asp Leu Glu Phe Pro 325 330 335

Ala Phe Thr Glu Lys Ser Ile Glu Asp Arg Arg Lys Met Tyr Asn Gln

345

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Leu Arg Glu Lys Glu Ile Asn Tyr Ala Asn Ser Het Val Gly Ser Pro 360 Asp Tyr Net Ala Leu Glu Val Leu Glu Gly Lys Lys Tyr Asp Phe Thr 370 380 Val Asp Tyr Trp Ser Leu Gly Cys Met Leu Phe Glu Ser Leu Val Gly Tyr Thr Pro Phe Ser Gly Ser Ser Thr Asn Glu Thr Tyr Asp Asn Leu Arg Arg Trp Lys Gln Thr Leu Arg Arg Pro Arg Gln Ser Asp Gly Arg Ala Ala Phe Ser Asp Arg Thr Trp Asp Leu Ile Thr Arg Leu Ile Ala Asp Pro Ile Asn Arg Leu Arg Ser Phe Glu His Val Lys Arg Met Ser 455 Tyr Phe Ala Asp Ile Asn Phe Ser Thr Leu Arg Ser Met Ile Pro Pro 470 475 Phe Thr Pro Gln Leu Asp Ser Glu Thr Asp Ala Gly Tyr Phe Asp Asp Phe Thr Ser Glu Ala Asp Met Ala Lys Tyr Ala Asp Val Phe Lys Arg 505 Gln Asp Lys Leu Thr Ala Met Val Asp Asp Ser Ala Val Ser Ser Lys Leu Val Gly Phe Thr Phe Arg His Arg Asn Gly Lys Gln Gly Ser Ser Gly Ile Leu Phe Asn Gly Leu Glu His Ser Asp Pro Phe Ser Thr Phe 545 550 Tyr



International Application No: PCT/

MICROORGANISMS									
Optional Sheet in connection with the microorganism referred to on page 108, lines 1-20 of the description									
A. IDENTIFICATION OF DEPOSIT'									
Further deposits are identified on an additional sheet *									
Name of depositary institution '									
American Type Culture Collection									
Address of depositary institution (including postal code and country) *									
12301 Parkiawn Drive Rockville, MD 20852									
US									
Date of deposit * March 24, 1995 Accession Number * 69769									
B. ADDITIONAL INDICATIONS ' (tenve blank if not applicable). This information is continued on a separate attached sheet									
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C. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE ' () As printed by the state of									
C. DESIGNATED STATES FOR WHICH RIDGE THE MADE BELLEVILLE									
D. SEPARATE FURNISHING OF INDICATIONS * (leave blank if not applicable)									
The indications listed below will be submitted to the internetional Bureau leter * (Specify the general nature of the indications e.g., *Accession flumber of Deposit*)									
E. I This sheet was received with the International application when filed (to be checked by the receiving Office)									
(Authorized Officer)									
☐ The date of receipt (from the applicant) by the International Bureau *									
was									
(Authorized Officer)									

Form PCT/RO/134 (January 1981)



### WHAT IS CLAIMED IS:

- 1. A purified lats protein.
- The protein of claim 1 which is a human protein.
  - 3. The protein of claim 1 which is a D. melanogaster protein.

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- 4. The protein of claim 1 which is a mouse protein.
- 5. The protein of claim 1 which is a mammalian 15 protein.
  - 6. The protein of claim 2 which comprises the amino acid sequence substantially as set forth in SEQ ID NO:4.

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7. A purified protein encoded by a nucleic acid hybridizable to the lats DNA sequence in plasmid PBS(KS)-h-lats as deposited with the ATCC and assigned accession number 69769.

- 8. A purified protein encoded by a nucleic acid hybridizable to a DNA having a sequence consisting of the coding region of SEQ ID NO:7.
- 9. The protein of claim 2 which is encoded by plasmid pBS(KS)-h-lats as deposited with the ATCC and assigned accession number 69769.
- 10. A purified derivative or analog of the protein 35 of claim 1, which displays one or more functional activities of a lats protein.

- 11. The derivative or analog of claim 10 which is able to be bound by an antibody directed against a lats protein.
- 5 12. A purified fragment of a lats protein comprising a domain of the lats protein selected from the group consisting of a lats C-terminal domain 3 (LCD3), lats C-terminal domain 1 (LCD1), kinase domain, a kinase subdomain, lats flanking domain 10 (LFD), lats split domain 1 (LSD1), lats split domain 2 (LSD2), SH3-binding domain, and opa repeat domain.
  - A molecule comprising the fragment of claim

14. A protein comprising an amino acid sequence that has at least 60% identity to a domain of a lats protein, in which the percentage identity is determined over an amino acid sequence of identical size to the domain.

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15. A protein comprising an amino acid sequence that has at least 90% identity to a domain of a lats protein, in which the percentage identity is determined over an amino acid sequence of identical size to the domain.

- 16. The derivative or analog of claim 10, which inhibits proliferation of a cell.
- 17. A chimeric protein comprising a fragment of a 30 lats protein consisting of at least 6 amino acids fused via a covalent bond to an amino acid sequence of a second protein, in which the second protein is not a lats protein.
- 18. The chimeric protein of claim 17 in which the 35 fragment of a lats protein is a fragment capable of being bound by an anti-lats antibody.



- 19. The fragment of claim 12 which additionally lacks one or more domains of the lats protein.
- 20. An antibody which is capable of binding a lats 5 protein.
  - 21. The antibody of claim 20 which is monoclonal.
- 22. A molecule comprising a fragment of the 10 antibody of claim 21, which fragment is capable of binding a lats protein.
  - 23. An isolated nucleic acid comprising a nucleotide sequence encoding a lats protein.

- 24. The nucleic acid of claim 23 which is a DNA.
- 25. An isolated nucleic acid comprising a nucleotide sequence complementary to the nucleotide sequence20 of claim 23.
  - 26. The nucleic acid of claim 23 in which the lats protein is a human lats protein.
- 27. An isolated nucleic acid comprising the lats coding sequence contained in plasmid pBS(KS)-h-lats as deposited with the ATCC and assigned accession number 69769.
- 28. An isolated nucleic acid hybridizable to the 30 lats DNA sequence in plasmid pBS(KS)-h-lats as deposited with the ATCC and assigned accession number 69769.
- 29. An isolated nucleic acid hybridizable to a DNA having a sequence consisting of the coding region of35 SEQ ID NO:7.

- 30. An isolated nucleic acid comprising a fragment of a lats gene consisting of at least 8 nucleotides.
- 31. An isolated nucleic acid comprising a
  5 nucleotide sequence encoding a fragment of a lats protein that displays one or more functional activities of the lats protein.
- 32. An isolated nucleic acid comprising a

  10 nucleotide sequence encoding the chimeric protein of claim

  17.
- 33. An isolated nucleic acid comprising a nucleotide sequence encoding a protein, said protein 15 comprising the amino acid sequence of SEQ ID NO:4.
  - 34. An isolated nucleic acid comprising a nucleotide sequence encoding the fragment of claim 12.
- 35. An isolated nucleic acid comprising a nucleotide sequence encoding the protein of claim 14.
- 36. A recombinant cell containing the nucleic acid of claim 23, in which the nucleotide sequence encoding the25 lats protein is under the control of a promoter that is not a native lats gene promoter.
  - 37. A recombinant cell containing a nucleic acid vector comprising the nucleic acid of claim 26.
  - 38. A recombinant cell containing the nucleic acid of claim 34.
- 39. A recombinant cell containing the nucleic acid 35 of claim 35.



- 40. A method of producing a lats protein comprising growing a recombinant cell containing the nucleic acid of claim 23, in which the nucleotide sequence encoding the lats protein is under the control of a promoter that is not a native lats gene promoter, such that the encoded lats protein is expressed by the cell, and recovering the expressed lats protein.
- 41. A method of producing a lats protein

  10 comprising growing a recombinant cell containing a nucleic acid vector comprising the nucleic acid of claim 26 such that the encoded lats protein is expressed by the cell, and recovering the expressed lats protein.
- 42. A method of producing a lats fragment comprising growing a recombinant cell containing the nucleic acid of claim 34 such that the encoded lats fragment is expressed by the cell, and recovering the expressed lats fragment.

- 43. A method of producing a protein comprising a fragment of a lats protein, which method comprises growing a recombinant cell containing the nucleic acid of claim 35 such that the encoded protein is expressed by the cell, and 25 recovering the expressed protein.
  - 44. The product of the process of claim 40.
  - 45. The product of the process of claim 41.

- 46. The product of the process of claim 42.
- 47. The product of the process of claim 43.
- 48. A pharmaceutical composition comprising a therapeutically effective amount of a lats protein; and a pharmaceutically acceptable carrier.

- 49. The composition of claim 48 in which the lats protein is a human lats protein.
- 50. A pharmaceutical composition comprising a 5 therapeutically effective amount of the fragment of claim 12; and a pharmaceutically acceptable carrier.
- 51. A pharmaceutical composition comprising a therapeutically effective amount of the protein of claim 14;10 and a pharmaceutically acceptable carrier.
  - 52. A pharmaceutical composition comprising a therapeutically effective amount of the chimeric protein of claim 17; and a pharmaceutically acceptable carrier.
  - 53. A pharmaceutical composition comprising a therapeutically effective amount of the nucleic acid of claim 23; and a pharmaceutically acceptable carrier.
- 20 54. A pharmaceutical composition comprising a therapeutically effective amount of the nucleic acid of claim 35; and a pharmaceutically acceptable carrier.
- 55. A pharmaceutical composition comprising a
  25 therapeutically effective amount of the recombinant cell of claim 36; and a pharmaceutically acceptable carrier.
- 56. A pharmaceutical composition comprising a therapeutically effective amount of an antibody that 30 immunospecifically binds to a lats protein; and a pharmaceutically acceptable carrier.
- 57. A pharmaceutical composition comprising a therapeutically effective amount of a fragment or derivative 35 of an antibody that immunospecifically binds to a lats protein, said fragment or derivative containing the binding



domain of the antibody; and a pharmaceutically acceptable carrier.

- 58. A method of treating or preventing a disease

  5 or disorder involving cell overproliferation in a subject
  comprising administering to a subject in which such treatment
  or prevention is desired a therapeutically effective amount
  of a molecule that promotes lats function.
- 59. The method according to claim 58 in which the disease or disorder is a malignancy.
- 60. The method according to claim 59 in which the disease or disorder is selected from the group consisting of 15 bladder cancer, breast cancer, colon cancer, leukemia, lung cancer, melanoma, pancreatic cancer, sarcoma, and uterine cancer.
- 61. The method according to claim 58 in which the 20 subject is a human.
- 62. The method according to claim 58 in which the disease or disorder is selected from the group consisting of premalignant conditions, benign tumors, hyperproliferative disorders, and benign dysproliferative disorders.
- 63. The method according to claim 58 in which the molecule that promotes lats function is selected from the group consisting of a lats protein, a lats derivative or analog that is active in inhibiting cell proliferation, a nucleic acid encoding a lats protein, and a nucleic acid encoding a lats derivative or analog that is active in inhibiting cell proliferation.
- 35 64. The method according to claim 58 in which the molecule that promotes lats function is a lats derivative or

analog that comprises a kinase domain of a lats protein that has been mutated so as to be dominantly active.

- 55. The method according to claim 58 in which the5 molecule that promotes lats function is the protein of claim14.
- or disorder involving a deficiency in cell proliferation or in which cell proliferation is desirable for treatment or prevention in a subject comprising administering to a subject in which such treatment or prevention is desired a therapeutically effective amount of a molecule that inhibits lats function.

- molecule that inhibits lats function is selected from the group consisting of an anti-lats antibody or a fragment or derivative thereof containing the binding region thereof, a lats derivative or analog that is capable of being bound by an anti-lats antibody and that is a dominant-negative protein kinase, a lats antisense nucleic acid, and a nucleic acid comprising at least a portion of a lats gene into which a heterologous nucleotide sequence has been inserted such that said heterologous sequence inactivates the biological activity of the at least a portion of the lats gene, in which the lats gene portion flanks the heterologous sequence so as to promote homologous recombination with a genomic lats gene.
- 30
  68. The method according to claim 66 in which the molecule that inhibits lats function is an oligonucleotide which (a) consists of at least six nucleotides; (b) comprises a sequence complementary to at least a portion of an RNA transcript of a lats gene; and (c) is hybridizable to the RNA transcript under moderately stringent conditions.

- 69. The method according to claim 66 in which the disease or disorder is selected from the group consisting of degenerative disorders, growth deficiencies, hypoproliferative disorders, physical trauma, lesions, and 5 wounds.
- 70. An isolated oligonucleotide consisting of at least six nucleotides, and comprising a sequence complementary to at least a portion of an RNA transcript of a lats gene, which oligonucleotide is hybridizable to the RNA transcript under moderately stringent conditions.
- 71. A pharmaceutical composition comprising the oligonucleotide of claim 70; and a pharmaceutically15 acceptable carrier.
- 72. A method of inhibiting the expression of a nucleic acid sequence encoding a lats protein in a cell comprising providing the cell with an effective amount of the oligonucleotide of claim 70.
- 73. A method of diagnosing a disease or disorder characterized by an aberrant level of lats RNA or protein in a subject, comprising measuring the level of lats RNA or protein in a sample derived from the subject, in which an increase or decrease in the level of lats RNA or protein, relative to the level of lats RNA or protein found in an analogous sample not having the disease or disorder indicates the presence of the disease or disorder in the subject.
- 74. A method of diagnosing or screening for the presence of or a predisposition for developing a disease or disorder involving cell overproliferation in a subject

  35 comprising measuring the level of lats protein, lats RNA or lats functional activity in a sample derived from the subject, in which a decrease in the level of lats protein,

lats RNA, or lats functional activity in the sample, relative to the level of lats protein, lats RNA, or lats functional activity found in an analogous sample not having the disease or disorder or a predisposition for developing the disease or 5 disorder, indicates the presence of the disease or disorder or a predisposition for developing the disease or disorder.

- 75. A method of diagnosing or screening for the presence of or a predisposition for developing a disease or 10 disorder involving cell overproliferation in a subject comprising detecting one or more mutations in lats DNA, RNA or protein derived from the subject in which the presence of said one or more mutations indicates the presence of the disease or disorder or a predisposition for developing the 15 disease or disorder.
- 76. A method of diagnosing or screening for the presence of or a predisposition for developing a disease or disorder involving a deficiency in cell proliferation or in 20 which cell proliferation is desirable for treatment or prevention in a subject comprising measuring the level of lats protein, lats RNA or lats functional activity in a sample derived from the subject, in which an increase in the level of lats protein, lats RNA, or lats functional activity in the sample, relative to the level of lats protein, lats RNA, or lats functional activity found in an analogous sample not having the disease or disorder or a predisposition for developing the disease or disorder, indicates the presence of the disease or disorder.
- 77. A kit comprising in one or more containers a molecule selected from the group consisting of an anti-lats antibody, a nucleic acid probe capable of hybridizing to a 35 lats RNA, or a pair of nucleic acid primers capable of priming amplification of at least a portion of a lats nucleic acid.

- 78. A kit comprising in a container a therapeutically effective amount of a lats protein.
- 79. A method of increasing cell growth in animals 5 or plants comprising inhibiting lats expression or activity in said animals or plants.
  - 80. The method of claim 79 in which cell growth is increased in an edible plant.

- 81. The method of claim 79 in which cell growth is increased in a farm animal.
- 82. A method of identifying a molecule that

  15 specifically binds to a ligand selected from the group consisting of a lats protein, a fragment of a lats protein comprising a domain of the protein, and a nucleic acid encoding the protein or fragment, comprising
  - (a) contacting said ligand with a plurality of molecules under conditions conducive to binding between said ligand and the molecules; and
  - (b) identifying a molecule within said plurality that specifically binds to said ligand.

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83. A recombinant non-human animal or plant that is the product of a process comprising introducing a nucleic acid encoding at least a domain of a lats protein into the plant or animal.

- 84. A recombinant plant containing and capable of expressing a *lats* antisense nucleic acid.
- 85. A recombinant non-human animal or plant in
  35 which a lats gene has been inactivated by a method comprising introducing a nucleic acid into the plant or animal or an ancestor thereof, which nucleic acid comprises a non-lats

sequence flanked by lats genomic sequences that promote homologous recombination.

- 86. A method of identifying a tumor suppressor
  5 gene comprising (a) identifying an overproliferation phenotype in a genetic mosaic; and (b) isolating a gene that is mutated in cells exhibiting said overproliferation phenotype.
- 10 87. The method of claim 86 in which the genetic mosaic is an animal containing (a) a nucleic acid encoding and capable of expressing a recombinase, and (b) intrachromosomal insertions of a target site at which the recombinase promotes recombination, on the homologous arms of both of a set of parental chromosomes; and the genetic mosaic has been produced by a method comprising inducing expression of the recombinase.
- 88. The method of claim 87 in which the
  20 recombinase is an FLP recombinase, and the target site is an
  FRT site.
- 89. The method according to claim 87 in which the recombinase is a Cre recombinase, and the target site is a 25 lox site.
  - 90. The method of claim 86 in which the overproliferation phenotype is the formation of overproliferated outgrowth tissue.

- 91. The method of claim 86 in which the overproliferation phenotype is the formation of a normal structure of larger than normal size.
- 92. A non-human mammal comprising (a) a nucleic acid sequence encoding a recombinase operably linked to a promoter; and (b) intrachromosomal insertions into the



homologous arms of both of a set of parental chromosomes, of a target site at which the recombinase can promote recombination.

- 5 93. The mammal of claim 92 which is heterozygous for an induced mutation.
- 94. The mammal of claim 93 in which the sequence encoding the recombinase is operably linked to an inducible 10 promoter.
  - 95. A method of making a genetic mosaic comprising inducing expression of the recombinase in the mammal of claim 93.

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- 96. A method for identifying a gene with an identifiable mutant phenotype comprising:
  - (a) identifying a mutant phenotype in a genetic mosaic animal, said genetic mosaic animal having been produced by a method comprising recombinantly expressing a recombinase within a cell of the animal so as to promote recombination at intrachromosomally inserted target sites on the homologous arms of both of a set of parental chromosomes; and
  - (b) isolating a gene that is mutated in cells exhibiting said mutant phenotype.
- 97. A method for identifying a gene with an 30 identifiable mutant phenotype comprising:
  - (a) identifying a mutant phenotype in a cultured cell, said cultured cell having been produced by a method comprising recombinantly expressing a recombinase within said cultured cell so as to promote recombination at intrachromosomally inserted target sites on

the homologous arms of both of a set of parental chromosomes; and

- (b) isolating a gene that is mutated in cells exhibiting said mutant phenotype.
- 98. The method of claim 97 in which the mutant phenotype is a transformed phenotype.
- 99. The mammal of claim 92 in which the promoter 10 is not a native recombinase gene promoter.
- 100. A method of inhibiting cellular senescence in a subject comprising administering to a subject in which such inhibition is desired an amount of a molecule that inhibits 15 lats function, effective to inhibit cellular senescence.
- 101. A method of inhibiting cellular senescence in cells in vitro comprising contacting cells in vitro with an amount of a molecule that inhibits lats function, effective 20 to inhibit cellular senescence.
- 102. The method according to claim 100 in which the molecule that inhibits lats function is selected from the group consisting of an anti-lats antibody or a fragment or
- 25 derivative thereof containing the binding region thereof, a lats derivative or analog that is capable of being bound by an anti-lats antibody and that is a dominant-negative protein kinase, a lats antisense nucleic acid, and a nucleic acid comprising at least a portion of a lats gene into which a
- 30 heterologous nucleotide sequence has been inserted such that said heterologous sequence inactivates the biological activity of the at least a portion of the lats gene, in which the lats gene portion flanks the heterologous sequence so as to promote homologous recombination with a genomic lats gene.
  - 103. The method according to claim 101 in which the molecule that inhibits lats function is selected from the

group consisting of an anti-lats antibody or a fragment or derivative thereof containing the binding region thereof, a lats derivative or analog that is capable of being bound by an anti-lats antibody and that is a dominant-negative protein kinase, a lats antisense nucleic acid, and a nucleic acid comprising at least a portion of a lats gene into which heterologous nucleotide sequence has been inserted such that said heterologous sequence inactivates the biological activity of the at least a portion of the lats gene, in which the lats gene portion flanks the heterologous sequence shas to promote homologous recombination with a genomic lats the

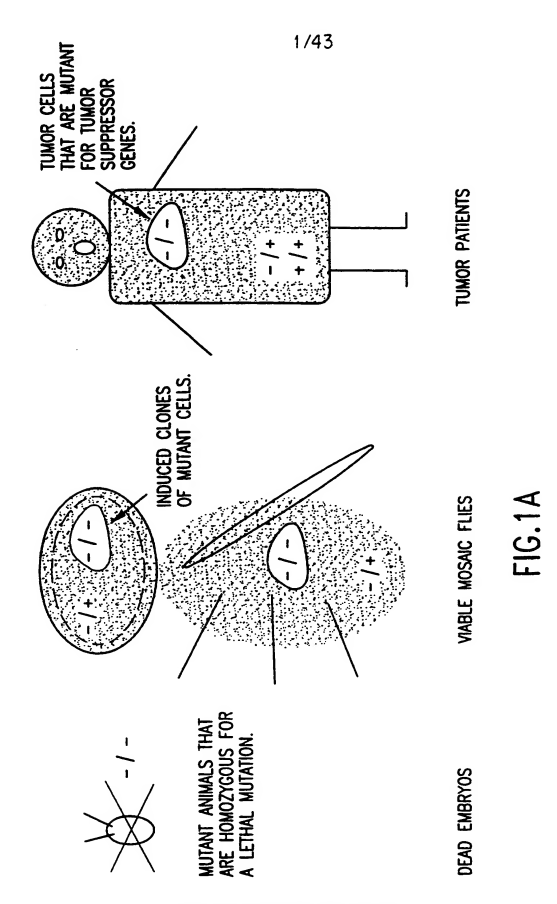
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**BAD ORIGINAL** 

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**SUBSTITUTE SHEET (RULE 26)** 

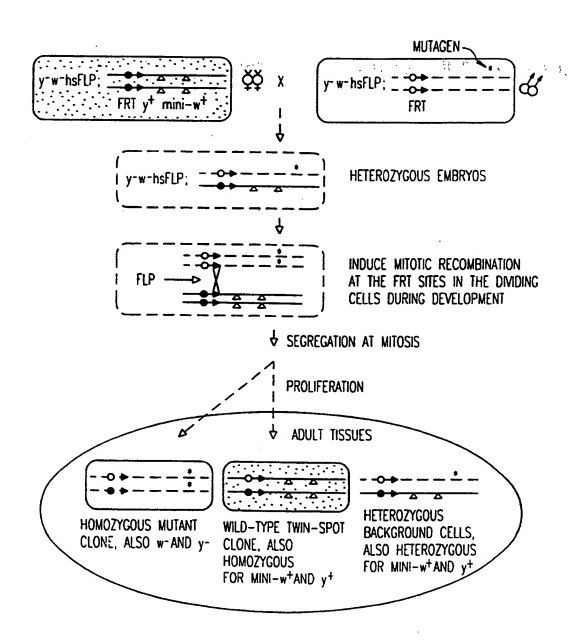


FIG.1B



FIG.2C

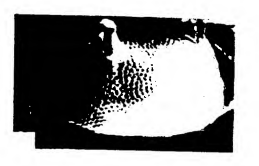


FIG.2B



FIG. 2,





FIG.2F



FIG.2E

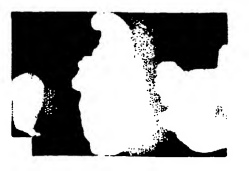


FIG.21



**FIG.2**I



FIG.2H



FIG.2G





FIG.2L

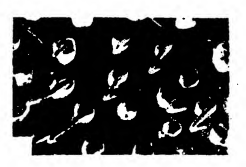
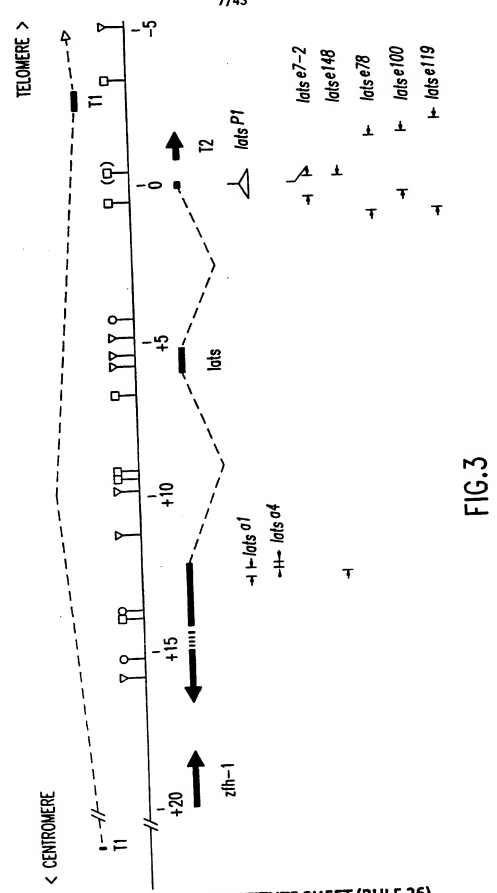


FIG.2K



FIG.2



SUBSTITUTE SHEET (RULE 26)

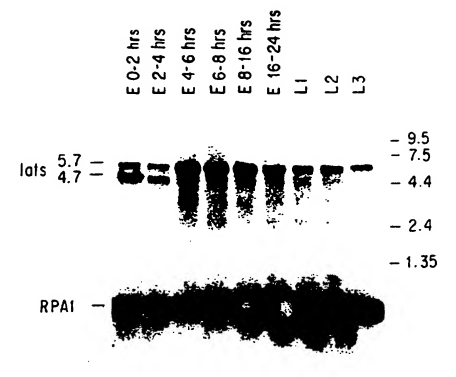


FIG.4

GENOA cDNA	AIC 1	9/43 ATCTAGCACGACGCAGCAACAAACCACGAATTAATTTTACTAAATTTAAGCCAAACGCGCATCGGAAATGCCT
	76	<u>A</u> GAAAATGCGATTGAATGCACGCGAAAAGTGATGCGTTGCGAACGCGACTGAATCAAGTGAAAATACGTCGGCAAA
	151	
	226	ACGCGAAAAAGGCGAGAACAAAGAGGCGAAAAGCGAGGAAATTGCGTGGAAAACGTGGAAAACGCGAAGAAGCGA
	220	INTRON 1 T
	301	AGCTCCAAGTTGGCCGCCATCGATTCGTGCGTAGGATCAATTAAGATTCCGAGTGGTCGAGAATCGGCTCAAATC
	376	AAATTAAAATCAACTAATATTTTGGTATTCAGATATTCAAATGGAATTCATTC
	451	ATCTGCCAACTATTTTTGAATTTGAATTGTGTGTCTGCGGCTGGCGCAGAATCTCTGATAAAGCAGAGGAATAAA
	526	ATCCGAAGAACAACAAATACAAATGAAATGAAATGCGCGGGGGGGG
	601	GGA GCGAAAĞGGGGGGTTTCTCTTATAATGCAAATGTGAATGTGAATGCGAATGCGAATGCGAGTGGAAGAATTCCCG
	676	GCGCGAGTGATAAATAATCCGACGACAAACAAAGCAGAAGCCTACACCGCGAGAAAGAGCAGCGCAAACACAATT G
	751	ATCTTTATTGAGAGCAACAATATCAAGATCGAGATAATAAAGCATCCTAAAACCCGCGCCTTAGTTCGTTTTAGT A
	826	CTCGCCACGGATATAGATATTCAAAGGCAAAAAGGTGGTGTCGGCATCGCCAGACAAACAA
	901	TCATACAAAACAACCAATTAAATAATAATAAAAAATAATA
	976	GCCGCCGATGTGCCCCCAGTGTGTGTGTGTGTGTGTGTGT
<b>A.</b> A.	1051	GAGCATTTCTGTGATATGAGTGCTAAATGCCACAGGGCGAAGCAGCAGCATCATGCATCCAGCGGGCGAAAAAAGG M H P A G E K R
	1126	GGGCGGTCGCCCAATGATAAATACACGGCGGAAGCCCTCGAGAGCATCAAGCAGGACCTAACCCGATTTGAAGT
	8	GGRPNDKYTAEALESIKQDLTRFEV INTRON 2
	1201	ACAAAATAACCATAGGAATAATCAGAATTACACACCTCTGCGATACACGGCGACCAACGGACGCAACGATGCACT
	34	Q N N H R N N Q N Y T P L R Y T A T N G R N D A L
	1276	TACTCCTGACTATCACCACGCCAAGCAGCCGATGGAGCCGCCACCCTCCGCCTCTCCTGCTCCGGACGTGGTCAT
	59	TPDYHHAKQPMEPPPSASPAPDVVI
	1351	
	84	PPPPAIVGQPGAGSISVSGVGVGVV
	1426	
	109	G V A N G R V P K M M T A L M P N K L 1 R K P S :
	1501	CGAACGGGACACGGCGAGCAGTCACTACCTGCGCTGCAGTCCGGCTCTGGACTCCGGAGCCGGTAGCTCCCGATC
	134	ERDTASSHYLRCSPALDSGAGSSRS
	1576	
	159	D S P H S H H T H Q P S S R T V G N P G G N G G F

FIG.5A SUBSTITUTE SHEET (RULE 26)

**BAD ORIGINAL** 



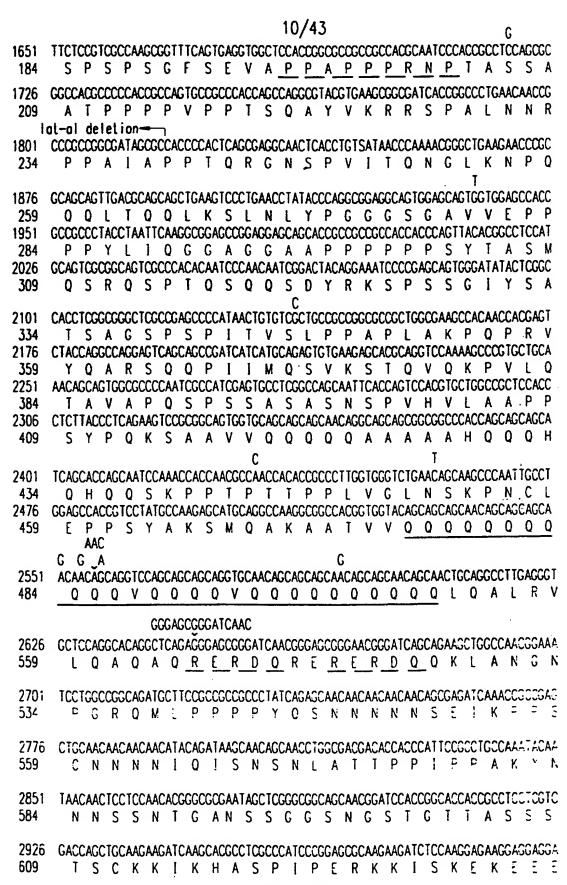


FIG.5B SUBSTITUTE SHEET (RULE 26)

**BAD ORIGINAL** 

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	11/43  1 GCGCAAGGAGTTCCGCATCAGGCAGTACTCGCCGCAAGCCTTCAAGTTCTTCATGGAGCAGCACATAGAGAACGT														_										
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3076 709	GATO				CGC R		CGC R	ACG T		CGC R		AAT N	CAG Q	CTG L	GAG. E	aag( K	GAG. E	ATG M	CAC. H	aaa K	GTG V	GGA G	CTG(	P	A D
3151 684	TCA	GACC T		ATC I	_		AGG R			CTG L		CAA			AGC S	AAC N	TAĊ Y	ATŤ l	CGA R	T TG L	AAG K	CGO R	GCC/ A	AAG/ K	T M
3226 759 3301	GGA D	CAA( K TAC(	ς	M	F	٧	K	L	K	Р	1	G	٧	G	Α	F	G	Ε	٧	T	L	٧	S	K	I
734	D		S			L	Y	A	M	K	T	L	R	K	A	D	٧	L	K	R	N	Q	V	. <b>A</b>	H
3376 809		gaai ' K			_	GAT D			XXX A		AGC(	D	N	)AA1 N ron	W	GTG V	GTC V	K K	TTC L	TAC Y	TAC Y	AGC S	TTC F	CAG Q	GA D
3451 784		GGA ( D					OTOT V			CTA( Y					IGA1 D			STCT	CTC	CTO	CATO I	AA/ K	ICTG L	GGC G	AT I
3526 809F	TTT	ICGA E	E	GGA E	L	GGC( A	CAG/ R	ATT( F	CTAI Y	CAT(	CGC A	CGA E	GGT( V	CAC( T	CTG( C	CCC A	OTOC V	GA( D	CAG( S	Ta V	CAC H	CAA/ K	M M	FFC G	TT F
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3676 859	GT(	GCAC C 1								CTC S	GAA K	GTA Y	CTA Y	CCA Q	GGA(	GAA! N	999 9	N	TCAI H tro	5	K	CCAI Q	GGAC D	S S	AT M
3751 884		•	PY	y E	E	Y	S	Ε	. 1	i G	F	) k	( F	·	V	L	Ē	R	R	K	М	A A	Ü	r.	Ĺ
3826 909	AA	GAG R	TCCT V (	TGG(	CCA	CTC ! S	GCT S L	GG1	GGC / C	CAC	CCC	GAA P N	\ \ -	CAT	AGC A	TCC	CGA E	GGT V	GCT L	GGA E	GAG R	GAG S	TGG: G	)A76 V	CAC
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3976 959	S CA	GTC S	CGC P	TGG/	AAA( E	000 <i>i</i> (	AACA ) (	<b>AA</b> ] [	AGG <sup>*</sup>	TCAT	CA i I	ACTO N I	GGG/ K	4GA/ E F	<b>VAAC</b> ( T	:CC	(30 <i>4</i> . H	[AT/	TCC	GCC	GCA C	(GCC ) 4	:03 <b>:</b> : <u>:</u>	377. -	A
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**SUBSTITUTE SHEET (RULE 26)** 

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1034	K H P T D T S N F D P V D P E K L R S N D S T M C C
4276	CAGCGCCGATGATGTCGATCAGAATGACCFCACTTTCCACGGCTTTTTCGAATTTACCTTCCGTCGCTTCTTCG
1059	SGDDVDQNDRTFHGFEFTFRRFF
4351	CGACAAGCAGCCCCCGATATGACGGACGATCAGGCCCCGTTTACGTCTGAAATGGATGCTCTCCATGTGCCC
1084	D, K Q P P D M T D D Q A P V Y V
4426	ACACCAACACCCCCCCCGAATCATTGTTAGTCAAATAGTCACAAAAAGGGGATAGAAACCATTGAGTGGGCT
4501	GCATTGTAAAGGAAGCCTGGGTATAGAATGAAACTATCTAT
4576	GGGAGCTACGTATATACATACAAATAATATACATATATTTGATATATAT
4651	ACTGAATAAATATAAAACGGAGCCGAGTAGAGATGAAACGAGAGGAGCGAGTCAGGACCTTCGACCTTTAACTG
4726	ACATAGTÁTATCCTTGTGCACTACTACTCCACAACAAATATATTTTTTAAATTGTTAGAATTCAAAAGGGACC
4801	ACTGGAAATCGAACCTTTCTGGTGCTCAAAGCAAAGCAA
4876	CGCGAATTTACCCAACCACTTCACTCCTCTCTCTCTCTCCACCTCCGATCGGTGGCCGGATTCGAACTCAGCAGG
4951	TGGTTGCATCCGGCCATCCCATTGACTTCCCATTCAGAATTGAGATTGCGAGGTGTGCGATGGAGAACGAAC
5026	GACCAAAAGTCGCACGCCAGCGATATAAGCCGGGTCTTATAAGCCTAATCTAAATCTAAACTGGGAGAACAGGAC
	C TGTAATTAGTG A A A
5101	TATGTATGTCCTGCTATCCAATTCGTCTATCACTGCTCTTCATCTGTGTACGACCCCCACCCCCCCC
5176	Identical to the 1-141 n.t. of the Drosophila pic-21 transcript CCAAAAGAACAACTIAGACGTAGCCTATGTGAAAAGCTAGCAATGTTAGACCAACTTGTTGAATGCCAAATGAA
5251	ATTGTTTAGCCCCATGAGGAAAACGCGGGGGAAATTCAACACTTATTCTCTGATAGCAAACGGAAAAGAAAG
5325	GAAAAAAAAAAAAAACAGAAACAGTACGAGAAAATTGTAATCTTCTTAATGTAATATTGTAAAFAACACFRRAARG
54["	AATSTATSSTAGAGTTGTGTGTAGCGCCCTAAGATGTTTTTTAGTTTATAGACCGCTAACCGTAATCTAGTTTAAT
547ê	COTAACACTAAGCGAGAGTACAGTACATTGGTTTTTTTGTTTG
5551	ACGATTIGTTTTTCTCTTTAATTAGCTTCAGTTTGTATGTGCGTGTGTTTTTATTATGACTTATATATA
5626	CTGAATATTCGTGGATGGAGCCTATTTTAAATGTGAGATCGAGCTAATTGAAGGAAATACAAACAA
5701	GCCTAGGCCAATTAGTTAT Poly A
	PAD ORIGINAL

FIG.5D

SUBSTITUTE SHEET (RULE 26)

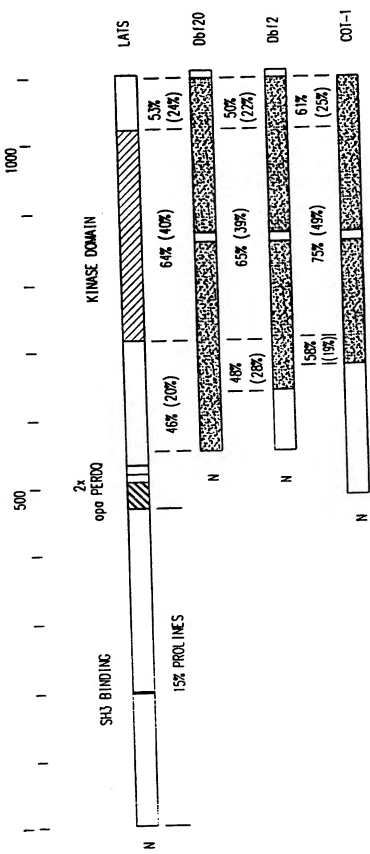


FIG.6A

SUBSTITUTE SHEET (RULE 26)



LATS DROSOPHILA 546 SINNNINGEJIKPPSGÅNINIOISNSØLATIPPIJFFANN-INISBNTGANSSÖLSNGBIGTTASSSTSOKKIIKHASFIIPERØKISKEKEFRIRETRIRENS PKIL7 TOBACCO   MCSARCEO   MCSARCON SOLVEN MCSARCON SOLVEN MCSARCON SOLVEN MCSARCON SOLVEN MCSARCON MCSARCON MCSARCON MCSARCON SOLVEN MEK	MFSRSDRÉINDDLAGÍÁNSHLGFYDLÍÐIR-KÐISÐGÁÐGRPARKÐENGRLIFÐLPRSYKPCDSDDQDIFÍÐARISLNHÐFIKKLPIÐ-DFHERASQSFIGÐVVNVC Eratsnætgevsvc	LATS DROSOPIILA 644 POÁFI-KIFMEDHIEMVIRSYRO-RTYRKNO-LEŘEMĚKVOLPDOTÓTEM ···RALINCKESNYIRLKRAKADKSMENALKATOVORFCEHVILVS-KIJÓTS COT-1 NEUROSPORA 191	ANY KOY IENGYME CAMIN-LOE PRE RRIL-LE MALADADVSE EDONIL LOGIL ENCETED MALICIA HAGADOE ELL TIMI CACAFOE PICMIGES VITTO	RKLEDADVSEEDONNL UKFLEHKKET KANKADOFFELLTMIOKSEGEHVR-IVC-REKTT	VIVIVA-FORDIENMENSOMKN-TOERKERRIW-LEKOLASSONPEERONSL INCLERKETTEFINRLKRANFICVNOFELLTTIII ORGANGEHVOTI-C-RFKKS	OLYTLOYYCOM-FOYVI-SRRO-RIKOVLRYLEDORSVKNYSNKVLNEEWALYLDAEHEVLAKARA KPKHKOFDILITOVAXXEKASHVALIAK-RAGILA	97 KMMFLEYYCOM-FOYVI-SRRQ-RIKOV-LEPLOQOSQLPNSDOJIKLNEEWSSYLOPEHOVLPKARALIOPKNROFEMITOVOOCKOLIMIJAR-IAOTI-	
1 546	— 83	644 191	43	-	4	<b>0</b>	97	
LAIS DROSOPHILA PKIL7 TOBACCO PK SPINACH	DBF 20 YEAST DBF 2 YEAST	LATS DROSOPHILA COT-1 NEUROSPORA	PK1L7 TOBACCO PK COMMON	ICE PLANT	PK SPINACH	<b>OBF 20 YEAS</b> 1	DBF2 YEAST	

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737 NII	259 GK	137 ONCRE		33 GH	. N3 96	193 DE	190 KE
1 737 NII LIYAWATERIADVEKRINDVAHVIKAERDILAEHDIAMINVIKLYYSFOOMONEYTIAMONIIPOGDUMBELINEGTFEFELIARFYJAEVICAVDSVHI	N 259 GK	137 ONCRE		63 CH	96 EN	193 DE	190 KE
IILA 737 NII	PORA 259 GK	.0 137 ONCRE		63 GH	96 EN	3T 193 DE	31 190 KE
OPHILA 737 NII	ROSPORA 259 GK	ACCO 137 ONCRE	Z.	63 GH	· N3 96 EN	EAST 193 DE	EAST 190 KE
ROSOPHILA 737 NII	NEUROSPORA 259 GK	TOBACCO 137 ONCRE	WOW	ANT 63 CH	INACH 96 EN	YEASI 193 DE	YEAST 190 KE
S DROSOPHILA 737 NII	-1 NEUROSPORA 259 GK	.7 TOBACCO 137 ONCRE	COMMON	PLANI 63 CH	SPINACH 96 EN	20 YEAST 193 DE	? YEASI 190 KE
ATS DROSOPHILA 737 NII	OT-1 NEUROSPORA 259 GK	KTL7 TOBACCO 137 ONCRE	IK COMMON	CE PLANT 63 GH	K SPINACH 96 EN	IBF 20 YEAST 193 DE	IBF2 YEASI 190 KE
LATS DROSOPHILA 737 NII	COI-1 NEUROSPORA 259 GK	PKTL7 TOBACCO 137 ONCRE	PK COMMON	ICE PLANT 63 GH	PK SPINACH 96 EN	DBF 20 YEAST 193 DE	OBF 2 YEAST 190 KE

FIG.6B

		•			
SPORA CO	DBI 2 YEAST  282 DLGYTHROLKPENFLIDAKHIKLTOFGLANGI 1 SNFRIESAKI -RLEKIKOLE -FPAFTEKSIEDR-RKMYNOL REKEINNAKARDDAMA  BBI 2 YEAST  282 DLGYTHROLKPENFLIDAKHIKLTOFGLANGI 1 SNFRIESAKI 1 SNFRIESAKI 1 SNFRIESAKI REKEINNAKARD KINASE DOMAIN KINASE DOMAIN KINASE DOMAIN KINASE DOMAIN KINASE DOMAIN RIGENSPORA 450 EIFIGINGYSFUCOWNSLIGHINFELLVOMPPFCMEDSHOWN INFRALL ON TE-RIGENCADELIKASHY FROYER PRITE TO DO THE SNOW SON THE SNICH		LAIS DROSOPHILA 1020 ADMAKOKÄPY IPEHKHYTÖTSÄFÖPÄDYEKLRÄNDSTÄKSÖDÜNDON COTI-NEUROSPORA 546 DSLIRRIRAPFEPRL ISAIDTITYFHPID-EHDOTONATLLKAODAARGAAAPAOOE PKIL7 TOBACCO 426 DRIYOMENAFIPENNDELIDTONFEKFE-ESESHSGSCSRSOPMRKML	ICE PLANT 344 LRIYOME AAF IPENDELIDIDNE KRE-EADNSSOSTSKÄDPMRKMLSKOLNENGYITYKNE IVNDYQVED IAELKKKIJIK + 55 A.A.  PR. SPINACII 385 INRI YE IDAAYKPONGELIDIDNE MKERIFEANPTPSRSOSOPSRKMLISKOLSE VGYITYKNEDAVKGLKHSF DRKGSTS + 39 A.A.  DBF 20 YEAST 475 ETTRISSPPF IPOLDOEITOÄGYF ODF TNEEDMAKYADVEKRONKL SAMV	

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* - I v.3vv.33	**	30	40 *	\$ * * * * * * * * * * * * * * * * * * *	60 *	70°2°2°2°2°2°2°2°2°2°2°2°2°2°2°2°2°2°2°2	80
	(1) II S I N R K (1) S W K G S K E S L V P Q R H G P S L V P Q R H	//////////////////////////////////////	1100/V/GG11 W K G 120	CIVAMBABIC S K E S 130	140 I V P 140 X X X X X X X X X X X X X X X X X X X	CAGNGNCACG Q R H 150	G P S L 160
CAAAAA I N 170	AGGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	GITCIGAAAG R S E S 190	CCCCAACTCA PNS 200	CAGGCGGATG Q A D 210	TAGGAAGACC V G R P 220	TCTGTCTGGA L S G 230	TCCGGCATTG S G I 240
2A11100 A 1 A 250 *	CAGCALLEGE FEAAGCTCAC CCAAGCAATG GACAGAGAGT GAACCCCCCA CCACCACCTC AAGTTAGGAG TGTTACTCCT A A 1 A 10 A 11 P S N G Q R V N P P P P Q V R S V T P 7.50	CCAAGCAATG P S N 270	GACAGAGAGT G Q R V 280	GAACCCCCA N P P 290	CCACCACCTC PPP 300	AAGTTAGGAG Q V R S 310	TGTTACTCCT V T P 320
CCACCTC.	CCACCACACA CACCCCACCT CCCCGAGGCA CCACTCCCCC TCCCCCCTCA TGGGAACCAA GCTCTCAGAC P P P P P S W E P S Q T 330 330 350 360 370 380 400	GACCCCACCT T P P 350	CCCCGAGGCA PRG 360	CCACTCCCCC T T P P 370	TCCCCCTCA P S 380	TGGGAACCAA W E P 390	GCTCTCAGAC S S Q T 400
AAAGCGCTAC TCTC K R Y S ATO	ICTGGGAACA S G N A20	IGGAGTACGT M E Y V 430	AATCICCCGA 1 S R 140	GNOCANCA IGGAGIACGT AATCICCCCG TTCCACCTGG GGCGTGGCAG GAGGGGTACC G N M Q E G Y C N M Q E G Y A N Q E G	TTCCACCTGG V P P G 460	GGCGTGGCAG A W Q 470	GAGGGGTACC E G Y 480
CTCCACCACC TCTT P P P P P P P P P P P P P P P P P P P	1 1 1 1 1 1 1 500 *	ICTCCCAIGA S P M 510	ATCCCCTAG N P P S 520	ACCAC:   ICTCCCA:   A TCCCCCTAG CCAGGCTCAG AGGCCCATTA GTTCTGITCC AGTTGGTAGA	AGGCCATTA R A I 540	GTTCTGTTCC S S V P 550	AGTTGGTAGA V G R 560
CCCATCA P 1	CANCCIALITA ICALIGOAGAG TACTAGCABA TITAACTITA CACCAGGGCG ACCTGGAGTT CAGAATGGTG GTGGTCAGTC	TACTAGCAAA T S K	TTTAACTITA F N I	CACCAGGGCG T P G R	ACCTGGAGTT P G V	CAGAATGGTG Q N G	GTGG1CAGTC G G Q S

640	TGATHITATIC GLIGGACCCAAAA ATGTCCCCAC TGGTTCTGTG ACTCGGCAGC CACCACCTCC ATATCCTCTG ACCCCAGCTA  10 F 1 V 11 Q N V P T G S V T R Q P P P Y P L T P A  11 CAULTUS CAUCAGCTA  12 CAUCAGCTA  13 CAUCAGCTA  14 CAUCAGCTA  15 CAUCAGCTA  16 CAUCAGCTA  16 CAUCAGCTA  17 CAUCAGCTA  18	A IGGACAAAG CCCCICTGCI TIACAAACAG GGGCTTCTGC TGCTCCACCA TCATTCGCCA ATGGAAACGT TCCTCAGTCG  N G () S P S A B P P S F A N G N V P Q S  730 740 750 760 770 780 790 800  * * * * * * * * * * * * * * * * * *	ATGATGGTOT: CCAACAGGAA CAGTCATAAC ATGGAGCTTT ATAATATTAA TGTCCCTGGA CTGCAAACAG CCTGGCCCCA M M M M M M M M M M M M M M M M M	CCAGTGAGGT PVR 960	AGTCACTGCC V T A 1040	ATCACACTOTO CHOCIALICA ACAGCCCGTG AAAAGCATGC GCGTCCTGAA ACCAGAGCTG CAGACTGCTY TAGCCCCAAC  1 1 1	AGTGTGCCTG S V P
630	ATATCCTCTG Y P L 710	ATGGAAACGT N G N V 790	CTGCAAACAG L Q T 870	ACCTAACATA P N I 950	CTGCCACTAC S A T T 1030	CAGACTGCTY Q 1 A 1110	TACAGCTTCA T A S
¢ *	CACCACCTCC P P P 700	TCATTCGCCA S F A 780	1GTCCCTGGA V P G 860	CTACATGGCA P T W Q 940	TCTCAGCCTT S Q P 1020	ACCAGAGCTG PEL 1100	TTTCTGAGGG F S E G
610	ACTCGGCAGC T R Q 690	TGCTCCACCA A P P 770	ATAATATTAA Y N I N 850	CATGAAATTC II E I 930	CTCTGCTAAT S A N 1010	GCGTCCTGAA R V L K 1090	CCTACCCCTT P I P
* 009	TGGTTCTGTG G S V 680	GGGCTTCTGC G A S A 760	ATGGAGCTTT M E L 840	AAGCGGTGGG S G G 920	GAGCAAGTCA R A S H 1000	AAAAGCA1GC K S M 1080	GACTGLICAG
\$ *	ATGTCCCCAC N V P T 670	TTACAAACAG L Q T 750	CAGTCATAAC S H N 830	AGTCA1CCCC () S S P 910	IINGGANGTA L G S 990	ACAGCCCGTG Q P V 1070	AGCCAGITCA
580	GIGCACCAAA V II Q 660 *	CCCCICTGCI P S A 740	CCAACAGGAA P N R N 820	000 V V V V V V V V V V V V V V V V V V	CAAATTCTTT TAATAACCTAC TAGGAAGTA GAGCAAGTCA CTCTGCTAAT TCTCAGCCTT CTGCCACTAC AGTCACTGCC S N S N S Q P S A T T V T A S N S Q P S A T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N T T V T A S N S N S N S N S N S N S N S N S N S	CICCIATICA A P 1 Q 1060	CCALCCLICIT ICALICCAC ACCCAGLICA GACIGLICAG CCTACCCCTT TTTCTGAGGG TACAGCTTCA AGTGTGCCTG  H P S H H P S H H P Q P V Q P V P F S E G T A S S V P
570	TGATITIALC D F 1 GEO	AFGGACAAAG N G Q S 730	ATGALGGICK M M V 810	GTCGTCTTCT S S S 890	CAAATICITI S N S L 970	ATCACACCCG 1 1 P 1050	CCATCCTTCT

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CGATACTAAA	CAAGAAAAGT	GTCTGTCTAG	GTTTGGTGAA	GAATAGGAGC	AGTOTATOTI TOTAMBATA ANGACATTAG GAATAGGAGC GTTTGGTGAA GTCTGTCTAG CAAGAAAAGT CGATACTAAA	IGIVAGAIA	AGTCTAIGLE
*	*	*	*	*	*	*	*
1680	1670	1660	1650	1640	1630	1620	1610
AAAATGGACA K M D	TAAAAGGGCT K R A	ATATTCGTCT Y I R L	GAGICIAACT E S N	TTGCCAGAAA C 0 K	AGATGCCCAG CATCAAATGA GAAAGATGCT TTGCCAGAAA GAGTCTAACT ATATTCGTCT TAAAAGGGCT AAAATGGACA 1) A () 1) Q M R K M L C Q K E S N Y I R L K R A K M D	CATCAMTGA  1) Q M	VGATGCCCAG D A Q
*	*	*	*	· *	*	*	*
6 L S Q	M R V	N E M	K Q L E 1570	H K K 1560	NVIENEMMRVGLSQ 1930 - 1540 - 1550 - 1560 - 1570 - 1580 - 1590 - 1600	0 H S 7 15/10	- N - N - N
GATTATCTCA	ATGCGGGTTG	AAATGAAATG	AGCAGCTAGA	CATCGGAAGA	GCAGCGTCTG	AGTCTCATCA	AACGICCICA
** 07CT	**************************************	*	*	* * * * * * * * * * * * * * * * * * *	) * [	*	; <del>;</del> -
H <	F M E Q	ъ ;	P 0 A	O S Y S	S R I	 ≃ ≃	K D C F
* GCACGTAGAG	* TCATGGAGCA	* TTTAAGTTCT	CCCACAGGCC	* AGAGTTACTC	* * * * * * * * * * * * * * * * * * *	ACGAAGAGAG	* AAGATGAAGA
1440	1430	1420	1410	1400	1390	1380	1370
N N	T V R	T S P I	0 I T	Ж	S G D K	S G D	S A D
* * * * * * * * GAGTGGGGAY, AGTGGTGACT CTGGGGATAA AGAAAAGAAA CAGATTACAA CTTCACCTAT CACTGTTGGG AAAAAGAAGA	* CACTGTTCGG	* CTTCACCTAT	* CAGATTACAA	* AGAAAAGAAA	* CTGGGGATAA	* AGTGGTGACT	* GAGTGCGGAC
1360	1350	1340	1330	1320	1310	1300	1290
D S E K	K E D	SLP	D E Q P	Р С К	V S K	PYES	S V
ATAGTGAGAA	AAGGAAGATG	TAGCTTACCC	ATGAACAGCC	CCCTGCAAAG	ICIGICCCIC CATATGAGIC AGTAAGTAAG CCCTGCAAAG ATGAACAGCC TAGCTTACCC AAGGAAGAIG ATAGIGAGAA	CATATGAGTC	1016100010
1280	1270	1260	1250	1240	1230	1220	1,210
CCAAAACCCA Q N P	ATCTGCTACA H L L H	TATCCAAAAC Y P K	ACCACCGCCT P P P	ATCAAGGTCC Y Q G P	TCATCCCACC IGITGCTGAA GCTCCAAGCT ATCAAGGTCC ACCACCGCCT TATCCAAAAC ATCTGCTACA CCAAAACCCA	IGITGCTGAA V A E	TCATCCCACC
*	**	*	0/17	*	OCT T	O+, T T	**************************************
1200	1190	1180	1170	1160	1150	1140	1130

1760	CGGAGAGGGA A E R D 1840 *	TACTTTGTGA Y F V 1920 *	ACGATTCTAC R F Y 2000	ATAGCAGAAC HACCIGIGC AGTTGAAAGT GTTCATAAAA TGGGTTTTAT TCATAGAGAT ATTAAACCTG ATAACATTTT  1 A L	GATIGACKEL CARCCAIA ITAAATIGAC TGACITTGGC TTGTGCACTG GCTTCAGATG GACACATGAC TCCAAGTACT  1 D R D G H I K L T D F G L C T G F R W T H D S K Y  1 D R D G H I K L T D F G L C T G F R W T H D S K Y  2100 2110 2120 2130 2140 2150 2160	ACCAGAGIGG GGAI CACCACACA GCAIGGATTI CAGTAACGAA TGGGGAGATC CTICCAATTG TCGGTGTGGG         Y       Q       S       G       D       S       M       D       F       S       N       C       G         Y       Q       S       G       D       S       M       D       F       S       N       C       R       C       G         Y       Q       S       G       D       S       M       D       F       S       N       C       C       G         P       S       G       D       S       M       D       F       S       N       C       C       G         P       S       G       D       S       M       D       F       S       N       C       N       C       G <th>GACAGACHGA ARCCACTGGA GCGGAGAGCT GCTCGCCAGC ACCAGCGATG TCTAGCCCAT TCTC1GGTTG GGACTCCCAA</th>	GACAGACHGA ARCCACTGGA GCGGAGAGCT GCTCGCCAGC ACCAGCGATG TCTAGCCCAT TCTC1GGTTG GGACTCCCAA
1750	CATGTGAAAG H V K 1830	GGACAACTTG D N L 1910	AAAATCTGGC E N L A 1990	ATTAAACCTG I K P 2070	GACACATGAC T H D 2150	CTTCCAATTG PSNC 2230	. TCTC1GGTTG S L V
1740	TCAGGTGGCT 0 V A 1820	TCCAGGACAA F Q D K 1900	ATCTTTCCTG I F P 1980	TCATAGAGAT H R D 2060	GCTTCAGATG G F R W 2140	TGGGGAGATC W G D 2220	TCTAGCCCAT
1730	TGCTCCGAAA L L R N 1810	TACTACTCTT Y Y S 1890	TAGAATGGGC R M G 1970	TGGGTTTTAT M G F I 2050	TTGTGCACTG L C T 2130	CAGTAACGAA S N E 2210	ACCAGCGATG H Q R C
1720	AAAGACGTTC K D V 1800	GGTCCGCCTG V R L 1880	GCCTATTAAT S L L I 1960	GTTCATAAAA V II K 2040	TGACITTGGC D F G 2120	GCATGGATTT S M D F 2200	GCTCGCCAGC A R 0
1710	TCTTCGAAAG L R K 1790	AIGAGTGGGT N E W V 1870	GATATGATGA D M M 1950	AGTTGAAAGT V E S 2030		CGGCAAGATA R Q D 2190	GCGGAGAGCT R R A
1700	CAACAAAGAC A T K 1 1780	CANGCCGACA F A D 1860	, 1CCTGGGGGG P G G 1940	11ACCTG1GC 1 1 C A 2020	CATGGCCATA D G H 2100	CGATCACCCA D H P 2180	AGCCACTGGA
1690	GCTTIGIAIG CAACAAAAGA AAAGACGTTC TGCTCCGAAA TCAGGTGGCT CATGTGAAAG CGGAGAGGGA A I. Y A T K I L R K K D V L L R N Q V A H V K A E R D A I. Y A T K I L R K K D V L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L L R N Q V A H V K A E R D N L R N Q V A E R D N L R N Q V A E R D N L R N Q V A E R N Q V A E R D N L R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N Q V A E R N	TATCCIAGCA CAAGCCGACA ATGAGTGGGT GGTCCGCCTG TACTACTCTT TCCAGGACAA GGACAACTTG TACTTTGTGA  I L A I A D N E W V V R L Y Y S F Q D K D N L Y F V  I L A I A D N E W V V R L Y Y S F Q D K D N L Y F V  I L A I A D N E W V V R L Y Y S F Q D K D N L Y F V  I R A I A D N E W V V R L Y Y S F Q D K D N L Y F V  I R A I A I A D N E W V V R L Y Y S F Q D K D N L Y F V  I R A I A I A I A I A I A I A I A I A I	TGGACTACAL ICCTGGGGGG GATATGATGA GCCTATTAAT TAGAATGGGC ATCTTTCCTG AAAATCTGGC ACGATTCTAC  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I F P E N L A R F Y  M D Y I P G G D M M S L L L I R M G I P F P E N L A R F Y  M D Y I P G G D M M M S L L L I R M G I P F P E N L R F Y  M D Y I P G G D M M M S L L L I R M G I P F P F P E N L R F Y  M D Y I P G G D M M M S L L L I R M G I P F P F F P F P F P F P F P F P F P F	1 A 1. 2010	6AT1GACCUT 1 0 R 1 0 R	ACCAGAG166 Y Q S G Y 170	GACAGACTOA D R I

		<b>~</b>			<b></b>								
2320	ATTCITTGTC I L C	2400	GCAAACTTC1	2480	CAGAAGACCG P E D R	2560	CTGAGACAGC L R Q 2640	*	TAAATTGTGG K L W	2720	AGCACGCTTT	2800	TATGAATACA Y E Y
2310	TGTTGGTGTT V G V	2390 *	TTATCATCTG	2470	TGTCGAGGAC C R G	2550 *	CTCTAGTGAT S S D 2630	*	TTGATCCTGA V D P D	2710	AAGCACCCCG K H P	2790	GCCTATTGAG P I F
2300	ACTGGTGGAG D W W S	2380	CAAATGAAGG	2960	TATCAAACTG I K L	2540	CCATCGATTT T I D F 2620	*	TTCGACCCTG F D P	2700	TAAAATGGG K N G	2780	ATTATCCAAA N Y P K
2290	CAGCTGTGTG Q L C	2370	ATTAGAAACA L E T	2450	CTGACCTCAT S D L I	2530	TTTTTTAGA F F K 2610	*	TACATCCAAT T S N	2690	GCGGATGGTA S G W Y	2770	TACCCATATA Y P Y
2280	AGGATATACA G Y T	2360 *	CACAAACCCC	2940	CCTGAAGCCT P E A	2520	GGCTCATCCA A H P 2600	*	ATCCAACAGA H P 1 D	2680	GACACTCIGA D T L	2760 *	TGACANTGGC D N G
2270	IACIGCGAAC I I R F	2350	CCTTCTTGG P F L	.2430 *	TAAGCTGAGT K L S	2510	AIGAGATAAA D E I K 2590	*	AAAATCACGC K I T	2670	AAATATCAGT N I S	2750 *	
2260	TEATATICICIA CICTICIAAGNIGIC TACTIGGGAACA AGGATATACA CAGCTGTGTG ACTGGTGGAG TGTTGGTGT ATTCTTTGTG  Y I A I' I V I I. R I G Y T Q L C D W W S V G V I L C	2340 *	AAAIGIIGGI GGGACAACCI CCIIICIIGG CACAAACCCC AITAGAAACA CAAATGAAGG TTAICAICTG GCAAACTICT E M L V G Q P P F L A Q T P L E T O M K V I I W O T S	2420	CIACACATORICO CINCLOANGO TAAGOTGAGT COTGAAGCOT CTGACCTCAT TATCAAACTG TGTCGAGGAC CAGAAGACCG	* *	AACGGTGCTG TL G A 2580	*	AIACAICCCI Y I P	* 0997	GCCACCACGCA	2740 *	ACHTEGGA F F R
2250	TFATATEGEA Y 1 A	2330 *	AAAIGTIGGI E M L V	2910	CTACACATCC L H I	0000	CCTCGGCAAG AACGGTGCTG ATGAGATAAA GGCTCATCCA TTTTTTAAGA CCATCGATTT CTCTAGTGAT CTGAGACAGC L C K T I D F S S D L R Q 25.70 25.80 25.90 26.00 26.10 26.20 26.30 26.40	₹:	AGICTÓCTIC ATACATOCO ANATOROGO ATOCAACAGA TACATOCAAT TTOGACCOTO TTGATOCTGA TAAATTGTGGO S A S Y I P K I T'H P I D T S N F D P V D P D K L W	2650 *	AGCGATGGTA GGGAGGGA AAATATCAGT GACACTCTGA GCGGATGGTA TAAAAATGGG AAGCACCCCG AGCACGCTTT S D T L S G W Y K N G K H P E H A F	2730	CIAIGMILL ACTUICGA GUITIIIGA FGACAATGGC TACCCATATA ATTATCCAAA GCCTATTGAG TATGAATACA Y I I I I II R P I I D D N G Y P Y N Y P K P I E Y E Y

9	ı	1	17
2	1	/	43

2880	ICALLICAGA GGGCICAGAA CAACAGICTG ATGAAGATGA TCAACACAA AGCTCCGATG GAAACAACCG AGATCTAGIG	TTGAGARAAT	3040	TTAAAATGTT 3120 *	ANTHIATHEE ACCITION TICAGIAATI TAGAAAAAT IGTTATAAGG AAAGTAAATT ATGAACTGAG TATTATAGIC 3200 3130 3180 3190 3200 3200 3130 4	INANGIN CITANAAAGA GANGCCIGGI AICTITIGIA TAIATAATAA AIAATTITAA AAICCCAAAA
2870	GAAACAACCG N N R 2950	TTTIGAAGTT	3030	GGAAATTGTT 3110 *	ATGAACTGAG 3190 *	ATATTTTAA
2860	AGCTCCGATG S S D G 2940	GTGCAGGGGT	3020	CTAAGTTATG G 3100	AAAGTAAATT A 3180 *	TATATAATAA
2850	TCAACACACA Q H T 2930	GAGGCCTGAA	3010	1111ATTTTC 3090 *	TGTTATAAGG A 3170	ATCTTTTGTA
2840	ATGAAGATGA D E D D 2920	GAA1 FTGCAA	3000	GTGTACAATA 3080	TAGAAAAAT 3160	GAAGCCTGGT
2830	CAACAGICTG Q Q S 2910	* ATCATTGTAA	2990	161616C1CT 3070	TTCAGTAATT 3150	CTTAMAAGA
2820	GGGCTCAGAA G S E 7900	* * * * * * * * * * * * * * * * * * *	2980	* * * * * * * * * * * * * * * * * * *	* ACCCITIIAA T 3140	
2810	7860 1 10 5 0 6 1 11 5 0 6	* * INIGITIMI	γ V γ 2970	* 1A1GCAAA1G 3050	* MITATICE / 3130	* ATTCHGGL ACT 3210

FIG. 7

WWWWWW WW

10	20 *	30	40	\$0 *	* 09	70	80
ATGAGAGCCA M R A 90	CCCCGAAGIT I P K F 100	IGGACCTIAT G P Y 110	CANAAAGCTC Q K A 120	ATGAGAGUCA CCCCGAAGGT IGGACCTIAT CAAAAAGCTC TCAGGGAAAT CCGATATTCC CTCCTGCCTT TTGCCAACGA M R A I I' K F G P Y Q K A L R E I R Y S L L P P A N E  90 100 110 120 130 140 150 160  * * * * * * * * * * * * * * * * * * *	CCGATATTCC R Y S 140	CTCCTGCCTT L L P 150	TTGCCAACGA P A N E 160
GTCAGGCAC1 S G 1 170	ICAGCAGCTG S A A 180	CAGAGGTGAA A E V N 190	CCGGCAGAIG R Q M 200	GTCAGGCACI ICGGCAGGTGAA CCGGCAGAIG CTTCAGGAGT TGGTGAATGC GGCATGTGAC CAGGAGATGG  S G I S A A A E V N R Q M L Q E L V N A A C D Q E M  170 180 190 200 210 220 230 240  * * * * * * * * * * * * * * * * * * *	TGGTGAATGC L V N A 220	GGCATGTGAC A C D 230	CAGGAGATGG Q E M 240
CTGGCAGAGC A G R A 250	GCTCACGCAG 1 1 0 260	ACGGCAG1A 1 G S 270	GGAGTATCGA R S I E 280	C1GGCAGAGC GTTATGA AGCTGCCTTG GAGTACATCA GTAAGATGGG CTACCTGGAC         A G R A I B R B I E A A L E Y I S K M G Y L D         250       270         280       300         310       320         320       310	GAGTACATCA E Y I 300	GTAAGA1GGG S K M G 310	CTACCTGGAC Y L D 320
CCCAGGAA1G P R N 330	ACCACA LIGHT 1 V 1 V 340	GCGAGICA1C R V 1 350	AAGCAGACCT K Q 1 360	CCAGGAATG AGTAGATCATC AAGCAGACCT CCCCAGGAAA GGGCCTGGCG TCCACCCCGG TGACTCGGCG  P R N 1 0 1 V R V 1 K Q 1 S P G K G L A S T P V T R R 330 330 330 350 360 370 380 390 400	GGGCCTGGCG G L A 380	TCCACCCGG S T P 390	TGACTCGGCG V T R R 400
GCCCAG111C P S F 410	GAGGGCACAG   G     A20	CCGAAGCACT G E A L 430	CCCATCCTAC P S Y 440	GCCCAGITIC GAGAGCACT CCCATCCTAC CACCAGCTGG GTGGTGCAAA CTACGAGGGC CCCGCCGCAC  P S F F G F A L P S Y H Q L G G A N Y E G P A A  A10	GTGGTGCAAA G G A N 460	CTACGAGGGC Y E G 470	CCCGCCGCAC P A A 480
1GGAGGAGA1 L E E M 490	GCCGCGCAA P R Q 500	IATI FAGACT Y L D: 510	TTCTCTTCCC F L F P 520	IGGAGGAGA   GCCCCCCA   IGGAGCCCCCA   ICT   ICT	GCCGGCACCC A G T 540	ACGGTGCCCA H G A Q 550	GGCTCACCAG A H Q 560
CATCCTCCTA H P P	AAGGGTACAG E. G. Y. S	CACAGCAGIA I A V	GAGCCAAGTG E P S	CATCCTCCTA ANGGRIACAG CATAGCAGIA GAGCCAAGTG CGCACTTTCC GGGCACACAC TATGGTCGTG GTCATCTACT II Y G R G II L L L L L	GGGCACACAC G T II	TATGGTCGTG Y G R	GTCATCTACT G II I. L

640	TCCAGCATGG S S M 720	CCAAGGCCCA GGGIGGCCCT CCGCCAGCC TCACCTITCC TGCCCATGCT GGGCTGTACA CTGCCTCGCA CCACAAGCCG  A K A Q G G P P A S L T F P A H A G L Y T A S H H K P  730 740 750 760 770 780 790 800	GGGGC CCACCCATTA CATGTGT1GG GCACCCGGGG TCCCACGTTT ACTGGCGAAA GCTCTGCACA   G A   I P L H V L G T R G P T F T G E S S A Q     B20	GGCTGTGCTG GCACCGGCCT CAATGCTGAC TTGTACGAGC TGGGCTCCAC GGTGCCCTGG TCTGCAGCTC  A V L A P S A A S L N A D L Y E L G S T V P W S A A A B B B B B B B B B B B B B B B B	CACTGGCACG CCGCAGACTCG AGGGTC1AGA AGCCTCGCGG CCGCATGTGG CTTT1CGGGC TGGCCCCAGC  P L A R R D S L Q K Q G L E A S R P H V A F R A G P S 8 P H V A F R A G P S 8 P H V A F R A G P S 8 P H V A F R A G P S 8 P S 990 1000 1010 1020 1030 1040 **	AGGACCAACI CCIICAACAA CCCACAACCT GAGCCCTCAC TGCCCGCCC CAACACGGTC ACCGCCGTGA CGGCCGCACAAAAAAAAAA	CALCOLLON: COUNTRANDA COGREGORIOT GOTGOGOCCO GAGOCCCAGA CAGOCGTGGG GCCCTCGCAC CCCGCCTGGG I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I II I V K S V R V L R P E P Q T A V G P S H P A W T I I II I V K S V R V L R P E P Q T A V G P S H P A W T I II I V K S V R V L R P E P Q T A V G P S H P A W T I I II I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I I V K S V R V L R P A W T I I I I I I V K S V R V L R P A W T I I I I I I V K S V R V L R P A W T I I I I I I V K S V R V L R P A W T I I I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I I I I V K S V R V L R P E P Q T A V G P S H P A W T I I I I I I I V K S V R V L R P R P M T I I I I I I I V K S V R V R V L R P R P T I I I I I I I I I I I V K S V R V R V R V R P R P T I I I I I I I I I I I I I I I I I I
630	AGATGCCTAT D A Y 710	CTGCCTCGCA T A S H 790	ACTGGCGAAA T G E 870	GGTGCCCTGG V P W 950	CTTT1CGGGC A F R A 1030	ACCGCCGTGA T A V 1110	GCCCTCGCAC P S H
620	AGACGCCACC K T P P X 700	GGGCTGTACA G L Y 780	TCCCACGTTT P T F 860	TGGGCTCCAC L G S T 940	CCGCATGTGG P H V 1020	CAACACGGTC N T V 1100	CAGCCGTGGG T A V G
610	TTCCAGAACA F Q N 690	TGCCCATGCT A H A 770	6CACCCGGGG G T R G 850	TTGTACGAGC L Y E 930	AGCCTCGCGG A S R 1010	TGCCCGCCC L P A P 1090	GAGCCCCAGA E P Q
009	CAGTTCCTCC S S S 680	TCACCTITCC L T F P 760	CATGTGT1GG H V L 840	CAATGCTGAC N A D 920	AGGGTC1AGA Q G L E 1000	GAGCCCTCAC E P S 1080	GCTGCGGCCC
290	66GTGCAGCG G V Q R 670	CCCGCCAGCC P A S 750	CCACCCATTA II P L 830	GGAACAGCCT R N S L 910	C1GCAGAAGC 1. Q K 990	CCCACAACCT P Q P 1070	CCGTGCGTGT S V R V
280	* 1C1GGGTATG S G Y 660	GGG1GGCCCT G G P 740	CACC1GGGGC P P G A 820	GCACCGICCA A P S 900	CCGCGAC1CG R D S 980	CCLICAACAAA S F N N 1060 *	CCIGIGAAGA P V K
570	ATCGGAGCAG 1C1GGGTATG GGGTGCAGCG CAGTTCCTCC TTCCAGAACA AGACGCCACC AGATGCCTAT TCCAGCATGG  S E Q S G Y G V Q R S S S F Q N K T P P D A Y S S M  S E Q S G Y G V Q R S S S F Q N K T P P D A Y S S M  650 670 680 690 700 710 720	CCAAGGCCCA A K A Q 730	GCGGCTACCC CACC16 A A I P P 810	GGCTGTGCTG A V L 890	CACTGGCACG P L A R 970	AGACCAACI R 1 N 1050	CALCCLICAC

1130	1140	1150	1160	1170	1180	1190	1200
TGGCTGCGCC V A A P LTTD	CACACCACCI 1 A P 1270	GCCACTGAGA A • T = E 1230	GCC1GGAGAC S L E T 1240	HANCHAURE CACACCCGCT GCTGGAGAC GAAGGAGGGC AGCGCAGGCC CACACCCGCT GGATGTGGAC	AGCGCAGGCC S A G 1260	CACACCCGCT PHPL 1270	GGATGTGGAC D V D 1280
(A1GGCGGC1) Y G G 1290	CCGAGCGCAG S F R R 1300	GTGCCCACCG C P P 1310	CCTCCG1ATC P Y 1320	TATGGCGGCCT	GCTGCTGCCC L L P 1340	AGTAAGTCTG S K S 1350	AGCAGTACAG E Q Y S 1360
CGTGGACCTG V D L 1370 *	CACAGCCTG1 D S L 1380	GCACCAGTGT C T S V 1390	GCAGCAGAGT Q Q S 1400	CGIGOACCIG CACCAGTGT GCACCAGAGT CTGCGAGGGG GCACTGATCT AGACGGGAGT GACAAGAGCC V D L D L D G S D K S V D L D L D G S D K S V D L D S L C T S V Q Q S L R G G T D L D G S D K S V D L D S L C T S V Q Q S L R G G T D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D G S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D K S V D L D C S D C	GCACTGATCT G T D L 1420	AGACGGGAGT D G S 1430	GACAAGAGCC D K S 1440
ACAAAGGTGC H K G A F450	GAAGGAGAC F G D F460	ANAGCTGGCA K A G 1470	GAGACAAAAA R D K K 11480	ACANAGOLOGO GAACAAAAAA GCAGATTCAG ACCTCCCGG TGCCTGTCCG CAAGAATAGC II K G I Q T S P V P V R K N S I K G I Q T S P V P V R K N S I K I I I I I I I I I I I I I I I I I	ACCTCCCCGG T S P 1500	TGCCTGTCCG V P V R 1510	CAAGAATAGC K N S 1520
AGAGATGAAG R D L 1530	AGAAGAGAGA 1 K. R. E. 1540 *	GICTCGCATC S R I 1550	AAGAG11AC1 K S Y 1560	AGABATICAAATTC TTCAIGGAGC AACACGTGGA R D L	CITCAAATTC F K F 1580	TTCA1GGAGC F M E 1590	AACACGTGGA Q H V E 1600
GAATGICATC N V I IG10	AAAACCTACC K 1 Y 1620	AGCAGAAGGT Q Q K V 1630	CAGCCGGAGG S R R 1640	GANTGLCALC ANANCCTACC AGCAGAAGGT CAGCCGGAGG CTACAGCTGG AGCAGGAAAT GGCCAAAGCT GGGCTCTGTG  N V   K   Y   Q   Q   K   V   S   R   L   Q   L   E   Q   E   M   A   K   A   G   L   C    16.10	AGCAGGAAAT E Q E M 1660	GGCCAAAGCT A K A 1670	GGCTCTGTG G L C 1680
AGGCGGAGGA F A 1 O	CCACCAGAIG	AGGAAGATCC R K I	TCTACCAGAA	ANTICITIANTIA CHANTIANTIA AGGAATEC TCTACCAGAA GGAGTCTAAC TACAACCGGC TGAAGAGGGC CAAGATGGAC F A L () L () M R K L L Y () K E S N Y N R L K R A K M D	TACAACCGGC Y N R	TGAAGAGGGC L K R A	CAAGATGGAC K M D

1690	1700	1710	1720	1730	1740	1750	1760
AAGICCATGI K S H 1770	ANGICCATGI HGIGAAAAT CAAGACTCTA GGCATCGGTG CCTTTGGGGA AGTGTGCCTC GCTTGTAAGC TGGACACTCA K S 11	CAAGACTCTA K T L 1790	GGCATCGGTG G I G 1800	CCTTTGGGGA A F G E 1810	AGTGTGCCTC V C L 1820	GCTTG1AAGC A C K 1830	TGGACACTCA L D T H 1840
CGCTCTGTAC A L Y 1850	CGCTCTGTAC GCATGAGA CTCTCAGGAA GAAGGATGTC CTGAACCGGA ATCAAGTGGC CCATGTCAAG GCTGAGAGGG  A L Y A M K T L R K K D V L N R N Q V A H V K A E R  1850 1860 1870 1880 1890 1900 1910 1920	CICTCAGGAA I L R K 1870	GAAGGATGTC K D V 1880	CTGAACCGGA L N R 1890	ATCAAGTGGC N Q V A 1900	CCATGTCAAG H V K 1910	GCTGAGAGGG A E R 1920
ACATCCTGGC D I I A 1930 *	ACALICETIGGE TGACAACT CTACTACTCC TTCCAGGACA AGGACAGCCT GTACTTTGTG  D I I A F A D N E W V V K L Y S F Q D K D S L Y F V  1930 1940 1950 1960 1970 1980 1990 2000  * * * * * * * * * * * * * * * * * *	AATGAGTGGG N E W 1950	TGGTCAAACT V V K L 1960	CTACTACTCC Y Y S 1970	TTCCAGGACA F Q D 1980	AGGACAGCCT K D S L 1990	GTACTTTGTG Y F V 2000
AIGGAC1ACA M D Y 2010	AIGGACTACA TATCAGGCGG GGATAIGATG AGCCTGCTGA TCAGGATGGA GGTCTTCCCT GAGCACCTGG CCCGCTTCTA M D Y 1 P G G D M M S L L I R M E V F P E H L A R F Y 2010 2020 2030 2090 2050 2060 2070 2080 * * * * * * * * * * * * * * * * * * *	GGATAIGATG D M M 2030	AGCCTGCTGA S L L 2090	TCAGGATGGA I R M E 2050	GGTCTTCCCT V F P 2060	GAGCACCTGG E H L 2070	CCCGCTTCTA A R F Y 2080
CALTGCAGAG I A L 2090	CATTGCACACAC TOTACCCACAAAG TGTCCACAAAG ATGGGCTTTA TCCACCGGGA CATCAAGCCT GACAACATAC I A I E S V II K M G F I H R D I K P D N I Z 1 V I K M G F I H R D I K P D N I Z 1 V I V M G F I H R D I K P D N I X M G F I H R D I K P D N I X M G F I H R D I K P D N I X I V I V I V I V I V I V I V I V I V	CCATGAAAG A I E S 2110	TGTCCACAAG V H K 2120	ATGGGCTTTA M G F 2130	TCCACCGGGA I H R D 2140	CATCAAGCCT I K P 2150	GACACATAC D N I 2160
1CATGGACCT L 1 D L 2170	1CALGGACCI GGAIGGICAT ALTAAGCIGA CAGATITIGG CCTCTGCACT GGATTCAGGT GGACTCACAA TTCCAAGTAC L D L D L D G H I K L T D F G L C T G F R W T H N S K Y 3.170 2180 2190 2200 2210 2220 2230 2240	AlfaAGC1GA 1 K L 2190	CAGATTTTGG T D F G 2200	CCTCTGCACT L C T 2210	GGATTCAGGT G F R 2220	GGACTCACAA W T H 2230	TTCCAAGTAC N S K Y 2240
TACCAGAAAG Y Q E	TACCAGACAAAAA GAAAAAAAAAAAAAAAAAAAAAAAAA	GAGACAGGAC R () D	AGCATGGAGC S M C	CCGGTGACCT P G D L	CTGGGACGAT W D D	GTTTCCAACT V S N	GTCGCTGTGG C R C G

o *	<b>\$</b> 0*		′∪ ⊢,	ນຸo∗	ຽ. 0.*	<u>∪.3.0</u> *	84 <del>1</del>
2320	ACACCA TP 240	TCTCTT L F 248	AGAGCA ES 256	GACTGC D C 264	CCGAAA R K 272	.GCCCCT S P 280	GCCTTC A F
C *	999 (	[ GA]	7 GGG	A A	CAI	E AAM	CAC =
2310	AGACAGGILA AMMACICITAS AGCAGAGGGC GCAGAAGCAG CACCAGAGGT GCCTGGCACA TTCTCTTGTC GGGACACCAA  D.R.L. K. I. L. F. Q. R. A. Q. H. Q. R. C. L. A. H. S. L. V. G. T. P.  2340 2350 2350 2350 2400  * * * * * * * * * * * * * * * * * *	ATTACATICAC TOUGGAGGIG CTTCTCCGCA AAGGGTACAC GCAGCTCTGT GACTGGTGGA GCGTCGGTGT GATTCTCTTT  N Y I A P E V L L R K G Y T Q L C D W W S V G V I L F  2410 2420 2430 2440 2450 2460 2470 2480  * * * * * * * * * * * * * * * * * * *	GAGATGCTGG THGGGCAGCC GCCTTTCTTG GCCCCCACCC CCACAGAGA GCAGCTGAAG GTGATCAACT GGGAGAGCAC  E M L V G Q P P F L A P T E T Q L K V I N W E S I  25.0 25.0 25.0 25.0 25.0 25.0 25.0 25.0	GCTGCATATC CCTACCACTCAG CGCTGAGGCC CGAGACCTCA TCACGAAGCT GTGCTGCGCG GCTGACTGCC L I T K L C C A A D C L L I T K L C C A A D C L R L L C C A A D C L R L L C C A A D C L R L L C C A A D C L R L L C C A A D C L R L L C C A A D C L R L L L C C A A D C L R L L L C C A A D C L R L L L C C A A D C L R L L L L C C A A D C L R L L L L L C C A A D C L R L L L L L C C A A D C L R L L L L L L L C C A A D C L R L L L L L L L L L L L L L L L L L	GUELIGAGGAAG   GUALIGAGGA CATCCGAAC   CATCCGAAC   CATCCGAAAG   CATCCAAAG   CATCCAAAAG   CATCCAAAAG   CATCCAAAAAA   CATCCAAAAAAAAAAAAAAAAAA	CAGGC 1 COLACG 1 CACCATCAGC CACCCCA IGG ACACCTCCAA TTTTGACCCG GTGGATGAAG AAAGCCCCTG Q A A P Y D E E S P W P T 1 S H P M D T S N F D P V D E E S P W P T 2730 2740 2750 2750 2770 2770 2780 2890 8800 *******************************	GCACGARRACO AGACCAAGGC CTGGGACACG CTGGCCTCCC CCAGCAGCAA GCATCCAGAG CACGCCTTCT II I A S P S S K H P E H A F
	TT S	S .	, GTG	<b>GT</b> 6	F	6T6 V	GCA
2300	GGCACA A H 2380 *	3GTGGA 4 W 2460 *	CTGAAG L K 2540	3AAGCT K L 2620	ICGACT I D 2700	3ACCCG D P 2780	CAGCAA S K
	GCCTI C L	GACT(	GCAG(	TCAC(	ACCA]	1111( F	CCAG(P S
2290 *	SAGGT R 2370	CTGT - C 2450	3AGAC E T 2530	CTCA L 2610	CAAC N 2690	S N 2770	STCCC S
	CACCA H Q	GCAGC Q I	CCACA(	CGAGA( R D	STTCT	ACACC:	CTGGC(
2280	6CAG 0 2360 *	ACAC Y T 2440	ACCC T	* V 7600 7600	ACCC (	AIGG / M   2760 *	CACG
	SCAGAA Q K	VAGGGT	SCCCC A P	GCTGA A E	GGCAC	ACCCC H P	TGGGA W D
2270	666C 6	GCA A R K 1430	L L 510	CAG C S 590	ICA A L K 670	. NGC C . S . 750	) (GGC (
	CAGAG	1CTCC	C1110	AGGCT R L	JGACC D	CCATC T I	CCCAN
<b>0</b> *	5 A 7 A 8 F	0 ×	ر د د عز د د عز	) ) ) ) ) ) )	\(\frac{1}{2}\)	\(\frac{1}{2} = 0 \cdot \)	ر ار ار
2260	CCCTG 1 L 234	GAGG1 E V 242	1057 10 120 120	CCCAG 1 0 758	0.00.00 0. A 266	CG1CC V 274	GAGAG G F
	₹	TCCC	) ) ) (	Z	CCV1	X	7.00 7.00 7.00 7.00 7.00 7.00 7.00 7.00
2250 *	% L 23330	VICGC 1 V 2410	C166 L 2490	VIAIC   1   255:0	300AG G R 2050 *	∩CAC A 2730 *	V()()()
	GACAI	TTTAC	AGA FO	C1GC/	CCTG(	אחמכ מ א	ICACG/
		Q Z	0		.5 &	C	9

2810 2820 2830 2840 2850 2860 2870 2880	
2860 2870  * CGAAGC CCTCAGAGCC P K P S E P 2940 2950  * STACGTG TAAGCCTCAG Y V * 3020 3030  * AATCCTT CATTTTTAGT 3100 3110  * TGCATTA AAACAGTATT	
2860 * CCGAAGC P K 2940 * 3020 AATCCTT AATCCTT AGCATTA	
TGCC C V AGG/	
2840 2850  \$\begin{array}{c} \cdot \	
2840 ACAACGGCTA D N G Y 2920 2920 A A E 3000 A CAGAAGACAA 3080 ATTAGCCCTC	. GGGGG
2830  11CTTCGATG A0  F F D D  2910  CTGGAAGGT G0  CATTTCTTTT C  3070  ATGATTTCAA A  ATGATTTCAA A	a CACITATIT
2820  * C11CCGCAGG 1 F R R 1 2900 * GGA1GCGA C G D A D 2980 * CCCAAAATGA G 3060 3140	TIAGIACACI ALGCAAAGAG CACITATITI GGGGG
2810  * ATGAGTICAC  Y E F 1  2870  * GCAGACCCAG  A D P  A D P  CTCGAGGAAA  3050  * GGGCAACAGG	* TIAGIACAG

FIG. 8F

08	AGCTCTGCTC 160	TTTCACTTTT 240	ATGAAGAGGA M K R	320	GTGAAAAGGC AGTAACTATA CTGTCAGTAG CCGGCAAATG S	TGAGTAAAAT M S K M 480	ATTCGAAACT I R N 560	AGACTTGCAA D L Q
70	6CTGTCCAGG 150 *	GAAGGATCAT 230 *	AGATGTTTTC	310	CTGTCAGTAG T V S S 390	GAGCATAACA E H N 470	CTTGCAGGAA L Q E 550	AAATGCTTCA Q M L Q
09	GCCCGTGGCC 140	GGACTTCCTT 220	CTACATATAT	300	AGTAACTATA S N Y 380	TGCTAAGGCT A K A 460	ATCATAAAGC H H K A 590	GTTAATCCAC V N P
50	CCGCCCTCAG 130	ACAGTCCTGG 210	TTCGTGTGGG	290	CTTTCCTGCC F P A 370	CATCTGATGC P S D A 450	1TTGGGACGC F G T 530	TACTTCAGAA T S E
40	GCCTCAGCGT 120	AAGAATTTTA 200 *	AAAGAAGTCC	280 *	GGCCTAAGAC R P K T 360	TTATCTAAAC L S K 440	TCCACCCAAA P P K 520	CTTCTCGGAG S S R S
30	\$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$	TIATATTGTA 190 *	ICTATCAAAT	270	AGACAAATGA R Q M 350	CCTTAGGAAT L R N 430	AAGTCAGAAA () V R N 510	GAAACAAATT E T N
20	IGCTGGGACG (	TCCCCTCCAG AGAGATITITA ACAGICCTGG GGACTICCTT GAAGGATCAT TTTCACTITT  170 180 190 200 210 220 230 240  * * * * * * * * * * * * * * * * * * *	GCTCAGAANA AANGELETGGA TETATCAAAT AAAGAAGTEE TTEGTGTGGG ETACATATAT AGATGTTTTE ATGAAGAGA M K R	\$ ×	AGAAGGATAT 1 G Y 340	TIACAAGAAA HEGGAAA CETTAGGAAT TTATCTAAAC CATCTGATGC TGCTAAGGCT GAGCATAACA TGAGTAAAAT  I. () I.   I.   S   I. R N L S K P S D A A K A E H N M S K M  A I ()   A 20 430 440 450 460 470 480  * * * * * * * * * * * * * * * * * * *	GICAACCGAA GATGGAAAA TCCACCCAAA TTTGGGACGC ATCATAAAGC CTTGCAGGAA ATTCGAAACT  S I F II P R () V R N P P K F G T H H K A L Q E I R N  A10 500 510 520 530 590 550 560  * * * * * * * * * * * * * * * * * * *	CTCTGCTTCC ATTTGCAAAT GAAACAAATF CTTCTCGGAG TACTTCAGAA GTTAATCCAC AAATGCTTCA AGACTTGCAA S L I P F A N E T N S S R S T S E V N P Q M L Q D L Q
10	ACC111GGG1 1GC1	TCCCCTCCAG	GCTCAGAAGA	**	GTGAAAAGCC S E K P 330	11ACAAGAAA 1. () 1 41()	GICAACCGAA S I E 490	CTCTGCTTCC S L I P

FIG. 9/

STO   SBO   SBO   SECTION   SECTIO								
S70   S80   S90   S90		CAATTGAATT A I E F 720	GCCAGCATGA A S M 800	TCAGAGGCAT Q R H 880 *	CTTTGTCTGG P L S G 960	CAAGTAAGGA Q V R 1040	ATGGGAACCA W E P 1120	GGGCATGGCA G A W Q
S70   S80   S90   600   610   620   8	e30 *	ATAGAAGCAG I E A 710	ACCTATTAAT P I N 790	CCTTAGTTCC S L V P 870	GTAGGAAGAC V G R 950	ACCACCACCT PPP 1030	CTCCCCTTC P P S 1110	GTCCCACCTG V P P
STO   SBO   SBO	620	CAACAGAAGT N R S 700	CAGCTGCCAG A A R 780	TCTAAAGAAT S K E 860	ACAGACAGAT Q T D 940	TGAACCCCCC V N P P 1020	ACAACTCCAC T T P 1100	AATCTCTCCT I S P
570   580   590   600	610	AGAAAACTAA Q K T N 690	ATGGCTGCAG M A A 770	CTGGAAAGGT W K G 850	GTCCCAACTC S P N S 930	GGACAGAGAG G Q R 1010	TCCAAGAGGT PRG 1090	TAATCTCCCG V I S R
GCIGCIGGAL HIGAIGAGGA TAIGGITATA  A A G L D E D M V I  GLO . 660 670  *  CALIAGIANA ALGAGTTACC AAGATCCTCG  I S K M S Y Q D P R  730 740 750  *  AACCAGGAAA IGIGCAGCAA TCAGITAAACC  K P G II V Q Q S V N  810 820 830  *  G P P I G E S V A Y  RPI G E S V A Y  AICHGIALA ICAGAGAAAAG IGIGGCCTAT  G P I G E S V A Y  810 900 910  *  AICHGIALA ICAGCALIIG ITCAAGCTCA  S G I S A I V Q A H  970 980 990  *  GIGITACTC ICCACCACCT CCAAGAGGCC  S V I P P P P P R G  1050 1050 1070  *  AACTCICAAA CAAAGCGCTA TCGGAAAC  N S O I K R Y S G N	009	CAAGCTCTTC Q A L 680	ACGAGAGCAG R E Q 760	GCAAACAGAG R K Q S 840	CATTCTGAGA H S E 920	CCCTAGCAAC P S N 1000	AGACTCCCCC Q T P P 1080	ATGGAATACG M E Y
570 580  * CCHGCIGGAL HGAIGAGGA  A A G I D E D G40 . 660  * CALIAGIAAA AIGAGTTACC I S K M S Y 730 740  * AACCAGGAA IGIGCAGCAA  K P G II V Q Q 810 820  * ACCCAGGAA IGIGCAGCAA  K P G II V Q Q 810 820  * AACCAGGAA IGIGCAGCAA  CGCCCACCAC I ACGAGAAAA  GGCCCACCAC I ACGAGAAAA  GGCCCACCAC I ACGAGAAAA  GGCCCACCAC I ACGAGAAAA  GIGITACHCI ICCACCACCI S V I P P P 1050 1060  * AACTCICAAA CAAAGCGCIA N S II K R Y	590	TAIGGF1ATA M V I 670	AAGATCCTCG Q D P R 750	TCAGTTAACC S V N 830	1GTGGCCTAT V A Y 910	1TCAAGCTCA V Q A H 990	CCAAGAGGCC P R G 1070	TTCTGGAAAC S G N
570 6C1GC1GGA1 A A G 640 * CATTAG1AAA I S K 730 * CATTAG1AAA K P G IJ 810 810 810 810 810 810 810 810	280		AIGAGTTACC M S Y 740	1G1GCAGCAA V Q Q 820	Iagagaaaga I G E S 900 *	1CAGCA111G S A F 980 *	ICCACCACCT P P P 1060	CAAAGCGCTA
	570	CCIGCIGGAL A A G A A G	CA11AG1AAA 1 S K 730	AACCAGGGAA K P G II 810	GGCCCGCCAC	A1CEGGIAIA S G 1 970 *	GIGITACICC S V I P 1050	MCTCTCAM N S 0

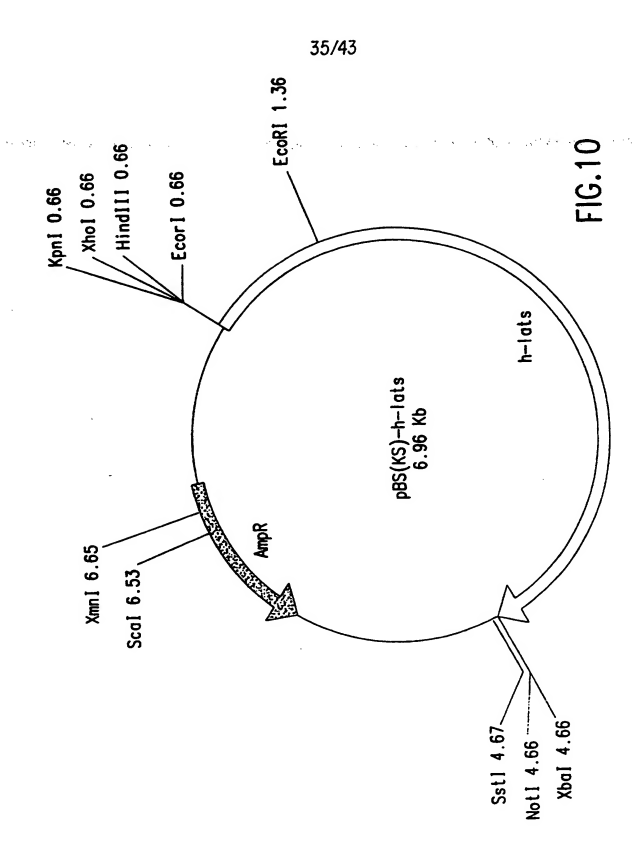
1190 1200	AGAGGGCIAT CCICCACCAC CTCTCACCATG AATCCTCCTA ATCAAGGACA GAGAGGCATT AGTTCTGTTC         E G Y P P P L N T S P M N P P N Q G Q R G I S S V         17:10       12:30         12:80       *         *       *	CTGTTGGCAG ACCAGCCAATC ATCAGCGA ATTTAACTTT CCATCAGGGA GACCTGGAAT GCAGAATGGT         P V G R Q P I I M Q S S K F N F P S G R P G M Q N G         1290       1300       1310       1320       1330       1350       1360         *       *       *       *       *       *       *	ACTGGACAAA CTGATTTCAT GATACACCAA AATGTTGTCC CTGCTGGCAC TGTGAATCGG CAGCCACCAC CTCCATATCC T G Q T D F M I H Q N V P A G T V N R Q P P P Y P 1370 1380 1390 1400 1410 1420 1430 1440 * * * * * * * * * * * * * * * * * * *	TCTGACAGCA GCTAATGGAC AAAGCCCTTC TGCTTTACAA ACAGGGGGGAT CTGCTGCTCC TTCGTCATAT ACAAATGGAA  L T A A N G Q S P S A L Q T G G S A A P S S Y T N G  1470 1480 1490 1500 1510 1520  * * * * * * * * * * * * * * * * * * *	GTATTCCTCA GTCTATGATG GTGCCAAACA GAAATAGTCA TAACATGGAA CTATATAACA TTAGTGTACC TGGACTGCAA S I P Q S M M V P N R N S H N M E L Y N I S V P G L Q 1530 1540 1550 1560 1570 1580 1590 1600 *	ACAAAIIGGC CTCAGTCATC TTCTGCTCCA GCCCAGTCAT CCCCGAGCAG TGGGCATGAA ATCCCTACAT GGCAACCTAA T N W P Q S S A P A Q S S G H E I P T W Q P N 1610 1620 1630 1640 1650 1660 1670 1680 * * * * * * * * * * * * * * * * * * *	CATACCAGIG AGGTCAAATT CTTTTAATAA CCCATTAGGA AATAGAGCAA GTCACTCTGC TAATTCTCAG CCTTCTGCTA
1180	ATCAAGGACA G N Q G Q 1260	CCATCAGGGA G P S G R 1340	TGTGAATCGG C V N R 1420	CTGCTGCTCC T S A A P 1500	CTATATAACA T L Y N I 1580	TGGCCATGAA A G H E 1660	GTCACTCTGC 1 S H S A
1170	AATCCTCCTA N P P 1250	ATTTAACTTT F N F 1330	CTGCTGGCAC P A G T 1410	ACAGGGGGAT T G G 1490	TAACATGGAA N M E 1570	CCCCGAGCAG S P S S 1650	AATAGAGCAA N R A
1160	TTCCCCCATG S P M 1240	GTTCTAGCAA S S S K 1320	AATGTTGTCC N V V 1400	TGCTTTACAA A L Q 1480	GAAATAGTCA R N S H 1560	GCCCAGTCAT A Q S 1640	CCCATTAGGA P I. G
1150	CTCTCAACAC P L N T 1230	ATCATGCAGA I M Q 1310	GATACACCAA I H Q 1390	AAAGCCCTTC Q S P S 1470	GTGCCAAACA V P N . 1550	TTCTGCTCCA S A P 1630	CTTTTAATAA S F N N
1140	CCTCCACCAC P P P 1220	ACAACCAATC Q P I 1300	CIGATTTCAT T D F M 1380	GCTAATGGAC A N G 1460	GTCTATGATG S M M 1540	CTCAGTCATC P Q S S 1620	AGGTCAAATT R S N
1130	AGAGGGCIAT E G Y 1210	CTGTTGGCAG P V G R 1290	ACTGGACAAA T G Q 1370	TCTGACAGCA L T A 1450	GTATTCCTCA S I P Q 1530	ACAAA11GGC T N W 1610	CATACCAGIG

1750 1760	CAACAGICAC IGCAAIIACA CCAGCTCCTA TTCAACAGCC TGTGAAAAGT ATGCGTGTAT TAAAACCAGA GCTACAGACT T V I A I P A P I Q Q P V K S M R V L K P E L Q T T V I A I P A P I Q Q P V K S M R V L K P E L Q T 17:0 17:0 17:0 17:0 18:0 18:0 18:0 18:0 18:0 18:0 18:0 18	ACACCACCC FICTTGGATA CCACAGCCAA TTCAAACTGT TCAACCCAGT CCTTITCCTG AGGGAACCGC  T II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P F P E G T A  1 II P S W I P Q P I Q T V Q P S P F P F P E G T A  1 II P S W I P C P S P F P F P F P F P F P F P F P F P F	TTCAAAIGIGA ACTATGC CACCTGTTGC TGAAGCTCCA AACTATCAAG GACCACCACC ACCTACCCA AAACATCTGC  S N V	TGCACCAAAA CECATICIGII CCTCCATACG AGTCAATCAG TAAGCCTAGC AAAGAGGATC AGCCAAGCTT GCCCAAGGAA  I II Q II P S V P P Y E S I S K P S K E D Q P S L P K E  I II Q II P S V P P Y E S I S K P S K E D Q P S L P K E  I II Q II P S V P P Y E S I S K P S K E D Q P S L P K E  I II Q II P S V P P Y E S I S K P S K E D Q P S L P K E  I II Q II P S V P P Y E S I S K P S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K P S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K P S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K E D Q P S L P K E  I III Q III P S V P P Y E S I S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K E D Q P S L P K E  I II Q III P S V P P Y E S I S K E D Q P S L P K E D Q P S L P K E D Q P S L P K E D Q P S L P K E P S K E D Q P S L P K E P S K E D Q P S L P K E P S K E D Q P S L P R E S L P R E S L P R E P S R E D Q P S L P R E S L P R E P S R E D Q P S L P R E S L P R E P S R E D Q P S L P R E S L P R E P S R E D Q P S L P R E S R E D Q P S L P R E S R E D Q P S L P R E S R E D Q P S L P R E S R E D Q P S L P R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E D Q P S R E S R E S R E D Q P S R E S R E D Q P S R E S R E S R E D Q P S R E S R E S R E S R E S R E S R E D Q P S R E	GAIGAGAGHT AAAACAGATT ACAACTTCAC CTATTACTGT  D F S   K S Y E N V D S G D K E K K Q I T T S P I T V  CHIQ 2100 2110 2120 2130 2140 2150 **	INGGNANANAC ANGANAGAGGANG GGAATCICGT ATTCAAAGTT ATTCTCCTCA AGCATTTAAA TTCTTTATGG R K N K K D E E R R E S R I Q S Y S P Q A F K F F M 21/0 2180 2290 2210 2220 2230 2240	AGCANCATGT MAAAAATGTA CTCAAATGC ATCAGCG TCTACATCGT AAAAAACAAT TAGAGAATGA AATGATGCGG
1740	ATGCGTGTAT T/ M R V L 1820	TCAACCCAGT CI 0 P S 1 1900	GACCACCACC A G P P P 1980 .	AAAGAGGATC A K E D Q 2060	GAAACAGATT A K Q I 2140	ATTCTCCTCA # Y S P Q 2220	AAAAAACAAT K
1730	TGTGAAAAGT V K S 1810	TTCAAACTGT I Q T V 1890	AACTATCAAG N Y Q 1970	TAAGCCTAGC K P S 2050	ATAAAGAAAA D K E K 2130	ATTCANAGTT I Q S 2210	TCTACATCGT
1720	TTCAACAGCC I Q Q P 1800	CCACAGCCAA P Q P 1880	TGAAGCTCCA E A P 1960	AGTCAATCAG E S 1 S 2010	GATAG1GGG D S G 2120	GGAATCICGT E S R 2200	ATCAGCAGCG
1710	CCAGCTCCTA P A P 1790	I I CTTGGATA S W I 1870	CACCTGTTGC P V A P 1950	CCTCCATACG P P Y 2030	IGAAAAIGIT L N V 2110 *	AAGAGCGAAG E E R R 2190	CICAAAICIC
1700	10CAA11ACA A 1 1 1780	CIACACACCC P I II P 1860	ACIGIGATGC 1 V M 1940	CCCALCIGIT P S V P S V P S 020	AAAAGAGIIA I K S Y 2100 *	AAGAAAGA 1G K K D 2180	V:3////////////////////////////////////
1690	CAACAGICAC T T V I	* GCTTTAGCAC CTAC A L A P 1 1850	TTCAAAIGIG S N V 1930	TGCACCAAAA I II Q II I	GATGAGAGTG D F S D P S	IAGGAAAAAC R K N 2170	VGCAACA161

	GIIGGAIIAI CICAAGAIGC CCAGGAICAA ATGAGAAAGA TGCTTTGCCA AAAAGAATCT AATTACATCC GTCTTAAAAG V G L S Q D A Q D Q M R K M L C Q K E S N Y I R L K R 2330 2340 2350 2350 2350 2360	GAAGTCTGT CTAGCAAGAA	2410 2420 2430 2440 2950 2460 2470 2480 * * * * * * * * * * * * * * * * * * *	ANGINGALAC INAGGETTIG TATGCAACAA AAACTETTCG AAAGAAAGAT GTTCTTCTTC GAAATCAAGT CGCTCATGTT  K V D I K A L Y A T K T L R K K D V L L R N Q V A H V  2490 2500 2510 2520 2530 2540 2550 2560	AAGGCTGAGA GACTGAAGCT GACAATGAAT GGGTAGTTCG TCTATATTAT TCATTCCAAG ATAAGGACAA K A I: R I) I I A E A D N E W V R L Y Y S F Q D K D N 7570 2580 2690 2600 2610 2620 2630 2640 * * * * * * * * * * * * * * * * * * *		TGGCACGALL CLACATAGCA GAACTTACCT GTGCAGTTGA AAGTGTTCAT AAAATGGGTT TTATTCATAG AGATATTAAA         1. A R I Y I A E L T C A V E S V H K M G F I H R D I K         27:30       27:40         27:30       27:50         27:30       27:50	CCTGAIAAIA IIIIGATTGA TCGTGATGGT CATATTAAAT TGACTGACTT TGGCCTCTGC ACTGGCTTCA GATGGACACA P II N I I I D R D G H I K L T D F G L C T G F R W T H
*	AAAAGAATCT / K E S 2380	GAGCATTTGG	2460 *	GTTCTTCTTC (	TCTATATTAT 1 L Y Y 2620	TAATTAGAAT G L I R M 2700	AAAATGGGTT 1 K M G F 2780	TGGCCTCTGC A G L C 1
*	TGCTTTGCCA M L C Q	CTAGGAATAG	2950 *	AAAGAAAGAT K K D 2530	GGGTAGTTCG W V R 2610	ATGAGCCTAT M S L 2690	AAGTGTTCAT S V H 2770	TGACTGACTT L T D F
*	ATGAGAAGA M R K	GATAAGACA	2440	AAACTCTTCG K T L R 2520	GACAATGAAT D N E 2600	GGGTGATA1G G D M 2680	GTGCAGTTGA C A V E 2760	CATATTAAAT H I K
*	CCAGGA1CAA Q D Q 2350	**************************************	2430	TATGCAACAA Y A T 2510	GGCTGAAGCT A E A 2590	ACATTCCTGG Y I P G 2670	GAACTTACCT E L T 2750	TCGTGATGGT R D G
*	CICAAGA1GC S Q D A	(ACAAGTCTA	2420	I/AGGCTTTG K A L 2500	GAGATATCCT R D I L 2580	CIAAIGGACI V M D 2660	C1ACA1AGCA Y 1 A 2740	IIIIGATTGA I I I D
*	G11GGA11A1 V G L 2330	\$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$	2410 *	AAGTAGATAC K V D T 2490 *	AAGGCTGAGA K A E 2570	111/1/1/11 1 Y 1 1 Y 1	TGGCACGAL1 L. A. R. I. 2730	CCTGATAATA P D N

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2880	2962	S 1	16TT( V 31;	11AT( / I 32(	75CCI	2700 S 334	TTGA V D
	A   C   S   G   D   S   M   D   F   S   N   E   M   G   D   S   S   S   D   D   S   S   S   S	GCIGICGAIG IGGAGACAGA CIGAAGCCAT TAGAGCGGAG AGCTGCACGC CAGCACCAGC GATGICTAGC ACATTCTTTG  S C R C G D R L K P L E R R A A R Q H Q R C L A H S L  2970 2980 3000 3010 3020 3030 3040  *	GTTGGGACTC CCAATTATAI TGCACCTGAA GTGTTGCTAC GAACAGGATA CACACAGTTG TGTGATTGGT GGAGTGTTGG  V G 1 P N Y 1 A P E V L L R T G Y T Q L C D W W S V G  3050 3060 3070 3080 3090 3100 3120  *	TGITATICH HIGAAAIGI TGGTGGGACA ACCTCCTTTC TTGGCACAAA CACCATTAGA AACACAAATG AAGGTTATCA  V I L I E M L V G Q P P F L A Q T P L E T Q M K V I  3130 3140 3150 3160 3170 3180 3200  * * * * * * * * * * * * * * * * * *	ACTGGCANAC ALCICLICAC ATTCCACCAC ANGCTAAACT CAGTCCTGAA GCTTCTGATC TTATTATTAA ACTTTGCCGA  N W Q I S L II I P P Q A K L S P E A S D L I I K L C R  3210 3220 3230 3240 3250 3260 3270 3280  * * * * * * * * * * * * * * * * * * *	GGACCCGANG AFCGCTTAGG CAAGAATGGT GCTGATGAAA TAAAAGCTCA TCCATTTTT AAAACAATTG ACTTCTCCAG G P E D R L G K N G A D E I K A H P F F K T I D F S S 3370 3380 3310 3320 3330 3340 3350 3360 *	TGACCIGAGA CAGCAGTOTG CTTCATACAT TCCTAAAATC ACACACCCAA CAGATACATC AAATTTTGAT CCTGTTGATC  1) IR () () S A S Y I P K I T H P T D T S N F D P V D
2870	666 (69 )	AGC A A 030	GGT W 1110	ATG M 1190	TAA K 8270 *	1 1 350 *	GAT
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2850	CATG M 293	CACG A R 301	GGAT G 309	ACAA 0 317	CTGA P E 325	6CTC A 333	4 P
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	7CCA P	TAG/	GTG] V	ACC.	ANG(	GCT(	700 P
2830	CCA (910)	CAT P 2990	FGAA E 3070	3ACA 3 0 3150 *	CCAC P 3230	1GGT G 3310	ACAT Y I
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AAAAGGAGAA	TGCTTTGTTT	<b>AAATAAGTAG</b>	<b>AAAGTACTTA</b>	CTTGGTACTT	<b>ATAGTCAGTT</b>	TANNITALGA ACTGAATATI ATAGTCAGTT CTTGGTACTT AAAGTACTTA AAATAAGTAG TGCTTTGTTT AAAAGGAGAA	TAMITAIGA
*	*	*	*	*	*	*	*
3920	3910	3900	3890	3880	3870	3860	3850
ATAAGGAAAG	<b>AAAAATTGTT</b>	<b>GTATTTAGAA</b>	GTTTAAATCA	IIIINA NNIGTTAATT TATTCCAGCC GTTTAAATCA GTATTTAGAA AAAAATTGTT ATAAGGAAAG	<b>AMIGITAATT</b>	AFCCITITA	ATTATEGGAA ATCCT
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3840	3830	3820	3810	3800	3790	3780	3770
ATTTTCCTAA	ACAATATTTT	TGCTCTGTGT	TATATATGTG	ATGACAGAGC	1 I ATGCAAAT	11GAGGIICI GAGAGIAAAA TIATGCAAAT ATGACAGAGC TATATATGTG TGCTCTGTGT ACAATATTTT ATTTTCCTAA	11GAGG11C1
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3760	3750	3740	3730	3720	3710	3700	3690
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GCGAGGTGTG	GGCCTGAAAT	TTTGTAAAG	GTAATGAGGA	GTAAATAAAT	TAACACACTA	AGIAIATGIT	ATCGCGATCT
3680	36/0	3660	3650	3640	3630	0.79s	9195
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GAGATTAAAA	CACAGGCTCA	ATGATCAAAA	TCGGATGAAG	AGAGCAGCAG	CACAAGGCTC	TGAATATGAA TACATTAATT CACAAGGCTC AGAGCAGCAG TCGGATGAAG ATGATCAAAA CACAGGCTCA GAGATTAAAA	TGAATATGAA
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CGAAGCCTAT	TATAATTATC	TGGCTACCCA	TTGATGACAA	CGAAGGTITT	ATTACCTIC	CALICIAIGA	CCIGAACAIG
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3520	3510	3500	3490	3480	3470	3460	3450
H Y 5	Z Y >	M C M	L O N	E E N <	D N E	N S D	р О Ж
TGGAAAGCAT	GGTATAAAAA	CTCAATGGAT	AAATGACACT	AAGAAATGT	GATAACGAGG	CTGATANALL ALGGAGTGAL GATAACGAGG AAGAAAATGT AAATGACACT CTCAATGGAT GGTATAAAAA TGGAAAGCAT	CTGATAAALI
*	*	*	*	*	*	*	*
3440	3430	3420	3410	3400	3390	3380	3370



**SUBSTITUTE SHEET (RULE 26)** 

EIVIS	III ATS PIKRSEKPEGYROMRPKTFPASNYTVSSROMLOETRESLRNLSKPSDAAKAEHNMSKMSTEDPRQVRNPPK 70
SIV III	HEATS FOILHERALOFIRNSLLPFANETNSSRSTSEVNPOMLQDLQAAGFDEDMVIQALQRINNRSIEAAIEFISK 140
EL AIS	III ATS PISYQDPRREQMAAAARPINASMKPGNVQQSVNRKQSWKGSKESLVPQRHGPPLGESVAYHSESPNSQTD 210 IIII ATS
E AIS	III ATS VCRPLSGSGTSAFVQAHPSNGQRVNPPPPPQVRSVTPPPPRGQTPPPRGTTPPPPSWEPNSQTKRYSGN 280
M.AIS m AIS	HLATS HEYVISRISPVPPGAWQEGYPPPPLNTSPMNPPNQGQRGISSVPVGRQPIIMQSSSRFNFPSGRPGMQNG 350 HILATS the state of the stat
ELAIS	III ATS TOQTOFMITIONVVPAGTVNRQPPPYPLTAANGQSPSALQTGGSAAPSSYTNGSIPQSMMVPNRNSHNME 420 IIII ATS 9.5.19t.s.tppap.fanv256
E A A A A A A A A A A A A A A A A A A A	HEALS EXPLISATED () INMPGSSSAPAQSSPSSGHEIPTWQPNIPVRSNSFNNPLGNRASHSANSQPSATTVTAIT 490 ml ATS
EL ALS	III ATS TAPTQQPVKSMRVLKPFLQTALAPTHPSWIPQPIQTVQPSPFPEGTASNVTVMPPVAEAPNYQGPPPPYP 560
III AIS	III ATS KIILLIIQNIYSVPPYESISKPSKEDQPSLPKEDESEKSYENVDSGDKEKKQITTSPITVRKNKKDEERRESR 630
III AIS	III ATS TOSYSPOATKFFMEQIVENVLKSHQQRLHRKKQLENEMMRVGLSQDAQDQMRKMLCQKESNYIRLKRAKM 700 mil ATS

1130 966	HEATS LIKNRIN VYV MLATS dan
1120	IN ATS DAILTEIVADEL NGWYKNGKIPEHAFYEFTFRRIFDDNGYPYNYPKPIEYEYINSQGSEQQSDEDDQNTGS 1120 MLATS GSissh.sh.s
1050 886	NLATS ASIN LIPLCRGPEDIRLGKNGADEIKAHPFFKTIDFSSDLRQQSASYIPKITHPTDTSNFDPVDPDKLWSD 1050
980 816	HLATS VGITNYTAPFVLLRTGYTQLCDWWSVGVILFEMLVGQPPFLAQTPLETQMKVINWQTSLHIPPQAKLSPE mLATS
910 746	INLATS HIKLIDFGLCTGFRWTHDSKYYQSGDHPRQDSMDFSNEWGDPSSCRCGDRLKPLERRAARQHQRCLAHSL MLATS
840 676	INLATS SEQUEDIN, YEVMDY FPGGDMMSLL IRMGTFPESLARFYTAEL TCAVESVHKMGFTHRD IKPONTL TORDG MLATS
909	TILATS DESPITED TO THE GLOVICAL VELAKKVOTRAL TATELLKENDY LLKINGVATIVAGENOTLAGADNEWY VILLT 7.00 ml ATS

=1G. 11B



hLATS mLATS2	hLATS MKRSEKPEGYRQMRPKTFPASNYTVSSRQMLQEIRESLRNLSKPSDAAKAEHNMSKMSTEDPRQVRNPPK mLATS2	70 45
hLATS mLATS2	III.AIS IGIIIIIKALQEIRNSLLPFANETNSSRSTSEVNPQMLQDLQAAGFDEDMVIQALQKTMNRSIEAAIEFISK nii AIS? pygooroonsgoossgoossgoossgoossgoossgoossgo	140 114
MLAIS MLAIS2	150 160 170 180 190 200 210 INLAIS MSYQDPRREQMAAAAARPINASMKPGNVQQSVNRKQSWKGSKESLVPQRHGPPLGESVAYHSESPNSQTD INLAIS2 g.l.n.i.vrvikqtspglastp.t.rp.fe.tg.asyqgang.aalee	210 175
HLA1S III A152	VGRPL.SGSG1SAFVQAHPSNGQRVNPPPPQVRSVTPPPPPRGQTPPPRGTTPPPPSWEPNSQTKRYSGN npj.qyldf1fpgag	280
INLATS INLATS?	NI YVISRISPVPPGAWQEGYPPPPLNISPMNPPNQGQRGISSVPVGRQPIIMQSSSKFNFPSGRPGMQNG adiliqaqahqhkstave.sahfpgthy.rghllseqsgygv.rs.q-nktp.dayss	350 251
H ATS mt ATS?	ld AIS - ICQIDI MIHQNVVPAGTVNRQPPPPPPL1AANGQSPSALQTGGSAAPSSYTNGSIPQSMMVPNRNSHNME mt AIS? տոեսգգգրթոslt.fpahaglytashhk-ptppgahp.hvl.trgtf.ge.sa.avla.sl.ad	420 319
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	BLATS TERRIN VYV 1130	14
1120	III ATS TEGES SAKAN-DII NGWYKNGKIIPETIAFYEFTI RRFF DONGYPYNYPKPIEYEYINSQGSEQQSDEDDQNTGS 1120 IIII ATS2 TEGESAKAN TEGES TO ASS TO A SECONDARY SAKAN	E VI
1050 948	HEATS ASPERTED TO THE CROPEDRICANGADE IKAHPEFKTIDESSDLRQQSASYIPKITHPTDTSNFDPVDPDICLWSD 1050 mil ATS? THE FLUCABORY. THE CABOR OF THE CA	E E
980	IN ALS VEHTNYTAPEVER RTGYTQLCDWWSVGVILFEMLVGQPPFLAQTPLETQMKVINWQTSLHIPPQAKLSPE 980 IN ALS?	HLAIS III AIS?
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770 668	HLATS DESHIVETEGIGAFGEVCLAREVDTKALYATETLREKDVLLRNQVAHVXAERDILAEADNEWVRLYY	IILATS IIILATS2
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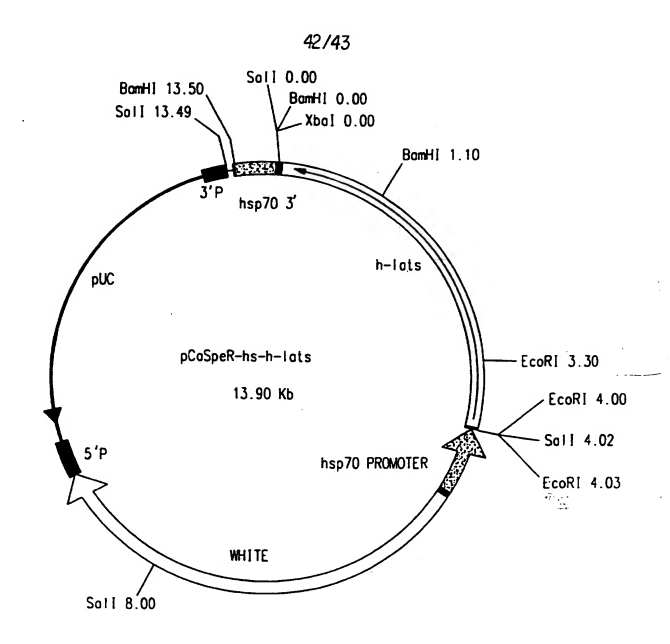


LSD20	
MKRSEKPEGYROMRPKTFPASNYTVSSROMLQEIRESLRNLSKPSDAAKAEHNMSKMSTEDPROVRNPPK-	70 30
FGTHHKALQEIRNSLLPFANETNSSRSTSEVNPOMLODLQAAGFDEDMVIQALOKTNNRSIEAAIEFISK	140
.evqnnhrnnqytp.rytagrndaltpdyhhakqpmepppsaspapdvv-ippppa.vgqpgag	97
MSYQDPRREQMAAAAARP I NASAKPGNVQQSVNRKQSWKGSKESLVPQRHGPPLGESVAYHSE-SPNSQTD	210
i.vsgvgvgvvgv.ngv-pmtalmpnklip.ierdta.shyl.cs.a.dsgagssrsdh.h-hSH3-BINDING	165
VGRPLSGSG1SAFVOAHPSNGQRVNPPPPPQVRSVTPPPPPRGQTPPPRGTTPPPPSWEPNSQTKRYSGN	280
thqs.rt.gnpggg-fs.s.sgfsevap.anp.assaa.pvpptsqayvr.pa	229
MEYVISRISPVPPGAWOEGYPPPPLNTSPMNPPNQGQRGISSVPVGRQPIIMOSSSKFNFPSGRPGMQNG	350
Innrppo.a.ptqrgnspvitqng.k-n.qqqlt.qlkslnly.g.gsgavveppppyliqg.ag.aapp	298
TGQTDFM1HQNVVPAGTVNRQPPPPYPLTAANGQSPSALQTGGSAAPSSYTNGS1PQSMAVPNRNSHNME	420
ppppsytasmqsrqsp.qsq.sd.rkspss.iytsaps.itvs1ppa.lakpq.rvyqarsq	364
LYNISVPCLQTNWPQSSSAPAQSSPSSCHEIPTWQPNIPVRSNSFNNPLGNRASHSANSQPSATTVTATT	490
qpi.mqsvks.qvqkpvlqtav.pqasasnspvhvlsappsypqksaavvqqqqqaaaaahqqqhqhq	436
LSD1a LSD2a LSD2p	
PAPIOOPVKSMRVLKPELQTALAPTHPSWIPQPIQTVQPSPFPEGTASNVTVMPPVAEAPNYQCPPPPYp	560
qskppt.ttpp1.glnskpnc.e.psyaksm.akaatvv erdqrererdqqklangnpgramlq	545
qqqqqqqqqqqqqqqqqqqqqqqqqqqqqqqqqqqqqq	
	MKRSEKPEGYROMRPKTFPASNYTVSSRCMLOE IRESLENL SKPSDAAKAEHNMSKMSTEDPROVRNPPK—  Mh.agekrggrpnd.yta.alesikqdltr  FGTHHKALOE IRNSLLPFANE TNSSRSTSEVNPOMLODLOAAGFDEDMVIQALOKTNNRSIEAAIEFISK .evqnnhrnnqytp.rytagrndaltpdyhhakqpmepppsaspapdvv-ippppa.vgqpgag  MSYQDPRREQMAAAAARPINASMKPGNVQQSVNRKQSWKGSKESLVPQRHCPPLGESVAYHSE-SPNSQTD i.vsgvgvgvvgv.ngv-pmtalmpnklip.ierdta.shyl.cs.a.dsgagssrsdh.h-h  SH3-BINDING  VGRPLSGSGISAFVQAHPSNGQRVNPPPPPDVRSVTPPPPPRGQTPPRGTTPPPPSWEPNSQTKRYSGN thqs.rt.gnpggg-is.s.sgisevapl.anp.assaa.pvpptsqayv.r.po  LSD1a  MEYVISRISPVPPCAWQEGYPPPPLNTSPMNPPNQCQRGISSVPVGRQPIIMQSSSKFNFPSGRPCMQNG Innrppa.a.ptqrgnspvitqng.k-n.qqqlt.qlksinly.g.gsgavveppppyliqg.ag.aopp  TGQTDFMIHQNVVPAGTVNRQPPPPYPLTAANGQSPSALQTGGSAAPSSYTNGSIPQSMMVPNRNSHNME ppppsytasmqsrqsp.qsq.sd.rkspss.iytsaps.itvslppa.lakpq.rvyqarsq  LYNISVPGLQTNMPQSSSAPAQSSPSSCHEIPTWQPNIPVRSNSFNNPLGNRASHSANSQPSATTVTAIT  qpi.mqsvks.qvqkpviqtav.pq  asasnspvhvlsoppsypqksaavvqqqqaaaaaahqqqhqhq LSD1a LSD1p LSD2a LSD2p  PAPIQQPVKSMRVLKPELQTALAPTHPSWIPQPIQTVQPSPFPEGTASNVTVMPPVAEAPNYQCPPPPYPp qskppt.ttppi.glnskpnc.e.psyaksm.akaatvv erdqrererdqqklangnpg\(\frac{qm1}{mm1}\)q

## FIG.13A

	41/43 LFD	
h-LATS LATS	KHLLHOMPSVPYESISKPSKEDQPSLPKEDESEKS-YENVDSGDKEKKQITTSPITVRKN—K-KDEERRESR qisnsnlattipvkynnnssntganssgg.ng.tgttas.stscikhapekis.e.ek.f.	
h-LATS LATS	IQSYSPQAFKFFMEQHVENVLKSHQQRLHRKKQLENEMMRVGLSQDAQDQMRKMLCQKESNYIRLKRAKM.rqi.i.yrtynkhkpdgt.ien	700 708
LF	TD KINASE DOMAIN	
h-LATS LATS	DKSMFVKIKTLGIGAFGEVCLARKVDT-KALYATKTLRKKDVLLRNQVAHVKAERDILAEADNEWVVRLYYpi.vl.vs.isnhmak	770 779
h-LATS LATS	SFQDKDNLYFVMDYIPGGDMMSLLIRMGIFPESLARFYIAELTCAVESVHKMGFIHRDIKPDNILIDRDG	840 849
h-LATS LATS	HIKLTDFGLCTGFRWTHDSKYYQ-SGDHPRQDSMDFSNEWGDPSSCRCGDRLKPLERRAARQHORCLAHSLnen.n.se-peey.e-npkptvrm.dv	910 915
h-LATS LATS	VGTPNYIAPEVLLRTGYTQLCKWWSVGVILFEMLVGQPPFLAQTPLETQMKVINWQTSLHIPPQAKLSPEe.syynsqekte.r.	980 985
	KINASE DOMAIN	
h–LATS LATS	.trrasadksvv.s.dga.m.k.k.pe.ker.n	1050 1053
h-LATS LATS		1120 1096
h-LATS LATS	EIKNRDUVYV 1130 1099	
	1003	

FIG.13B



**FIG.14** 

	FETAL	ADULT	
	BRAIN LUNG LIVER KIDNEY	SPLEEN THYMUS PROSTATE TESTIS OVARY SMALL INTESTINE	BLOOD LEUKOCYTE
h-lats			— 9.5 — 7.5 — 4.4
β-actin			— 2.4 — 1.35

FIG.15

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/04101

A. CL.	ASSIFICATION OF SUBJECT MATTER :C07K 11/00; C07H 21/04; C12P 21/02; C12N 5	5/10: A61K 38/43	
US CL According	:530/350; 536/23.2; 435/69.1, 240.1; 514/2 to International Patent Classification (IPC) or to b		·
	LDS SEARCHED		
Minimum o	documentation searched (classification system follo	wed by classification symbols)	
	530/350; 536/23.2, 23.4; 435/69.1, 69.7, 240.1;		
Documenta	tion searched other than minimum documentation to	the extent that such documents are include	ed in the fields searched
Electronic o	data base consulted during the international search	(name of data base and, where practicable	e, search terms used)
APS, BIC	OSIS, IntelliGenetics erms: lats gene, drosophila tumor suppressor g		
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.
X  Y	GENES AND DEVELOPMENT, Vo 01 March 1995, Justice et al Suppressor Gene warts Encode Myotonic Dystrophy Kinase and is Cell Shape and Proliferation", p document.	l, "The <i>Drosophila</i> Tumor es a Homolog of Human s Required for the Control of	19, 23-25, 28- 32, 34-36, 38- 39 
X Further	EMBO JOURNAL, Volume 11, Nu Yarden et al, "cot-1, a Gene Requin Neurospora crassa, Encodes 2159-2166, see entire document	uired for Hyphal Elongation a Protein Kinase", pages	19, 28-30, 32
	r documents are listed in the continuation of Box (		
A* docu	ial categories of cited documents: ment defining the general state of the art which is not considered of particular relevance	"T" later document published after the inte date and not in conflict with the applic principle or theory underlying the inv	tion but cited to understand the
docu	er document published on or after the international filing date ment which may throw doubts on priority claim(s) or which is to establish the publication date of another citation or other	"X" document of particular relevance; the considered novel or cannot be considered when the document is taken alone	e claimed invention cannot be red to involve an inventive step
specia	ne reason (as specified)  ment referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance; the considered to involve an inventive combined with one or more other such being obvious to a person skilled in the	step when the document is documents, such combination
docur the pr	nent published prior to the international filing date but later than nority date chained	*&* document member of the same patent	
ate of the ac	tual completion of the international search	Date of mailing of the international sea	rch report
Commissioner Box PCT Washington, I Icsimile No.	illing address of the ISA/US r of Patents and Trademarks D.C. 20231 (703) 305-3230 /210 (second sheet)(July 1992)#	Authorized officer  ROBERT C. HAYES  Telephone No. (703) 308-0196	tesde



#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/04101

		PC1/US96/041	01
C (Continu	ation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relev	ant passages	Relevant to claim No
x	THE JOURNAL OF BIOLOGICAL CHEMISTRY, Volume 266, Number 19, issued 05 July 1991, Shortridge et al, "A <i>Drosophila</i> Phospholipase C Gene that is Expressed in the Central Nervous System", pages 12474-12480, see entire document.		7-8, 10-11, 14- 15, 17-19, 28-30 32, 35, 39
X	GENE, Volume 104, Number 1, issued 1991, Toyn et Cell-Cycle-Regulated Budding Yeast Gene <i>DBF2</i> , Enco Putative Protein Kinase, has a Homologue that is Not Cycle Control*, pages 63-70, see entire document.	oding a	7-8, 10-11, 14- 15, 17-19, 28-30 32, 35, 39
X, P  Y, P	Xu et al, "Identifying Tumor Suppressors in Genetic Mosaics: the		1, 3, 6-8, 10-19, 23-25, 28-32, 34-36, 38-39
	·		2, 4-5, 9, 26-27, 33, 37, 40-52, 78



Intern anal application No. PCT/US96/04101

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)					
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:					
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:					
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:					
Claims Nos.:  because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).					
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)					
This International Searching Authority found multiple inventions in this international application, as follows:					
Please See Extra Sheet.					
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchal claims.	əlc				
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payme of any additional fee.	:nt				
3. As only some of the required additional search fees were timely paid by the applicant, this international search report cover only those claims for which fees were paid, specifically claims Nos.:	:rs.				
4. X No required additional search fees were timely paid by the applicant. Consequently, this international search report restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-19, 23-52, and 78	is				
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.					



#### INTERNATIONAL SEARCH REPORT



International application No. PCT/US96/04101

unity of invention is lacking.

Groups V and XIV contain claims directed to more than one species of the generic invention. These species are deemed to lack Unity of Invention for the above reasons, which explain why the compositions used lack unity and are not so linked as to form a single inventive concept under PCT Rule 13.1. If the fee for searching Groups V or XIV is paid, the first named embodiment the anti-lats antibody, will be searched. In order for more than one species to be examined, the appropriate additional examination fees must be paid. The species for claims 66-67, 69, 100-103 are as follows:

- A) anti-lats antibody.
- B) lats derivative or analog.
- C) lats antisense nucleic acid.
- D) a nucleic acid comprising a portion of the lats gene.

In Group V, the following claims are generic: claims 66-67, 69. In Group XIV, the following claims are generic: claims 100-103.

#### INTERNATIONAL SEARCH REPORT

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BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group I, claims 1-19, 23-52 and 78, drawn to a purified lats protein, derivative, analog, or fragment, a chimeric protein, an isolated nucleic acid, a recombinant cell, a method of producing the lats protein and a pharmaceutical composition and a kit that comprises a lats protein.

Group II, claims 20-22, 56-57 and 77, drawn to an antibody, a molecule comprising antibody fragments and a pharmaceutical composition and a kit comprising these antibodies/fragments.

Group III, claims 53-55, 70-71 and 77, drawn to pharmaceutical compositions comprising a therapeutic nucleic acid, an oligonucleotide, a recombinant cell and a kit comprising the nucleic acid probes/primers.

Group IV, claims 58-65, drawn to a method of treating a disease state by administrating a molecule that promotes lats function.

Group V. claims 66-69, drawn to a method of treating a disease state by administrating a molecule that inhibits lats function.

Group VI, claim 72, drawn to a method of inhibiting expression of a nucleic acid with an oligonucleotide.

Group VII, claims 73-76, drawn to a method of diagnosis of a disease by screening aberrant levels of lats RNA or protein using nucleic acids or proteins or antibodies.

Group VIII, claims 79-80, drawn to a method to increase cell growth in plants.

Group IX, claims 79 and 81, drawn to a method to increase cell growth in animals.

Group X, claim 82, drawn to a method of screening for lats ligands.

Group XI, claims 83-85, drawn to transgenic plants.

Group XII, claims 83, 85, 92-95 and 99, drawn to transgenic animals and method of making.

Group XIII, claims 86-91 and 96-98, drawn to a method of identifying a tumor suppressor gene.

Group XIV, claims 100-103, drawn to a method of inhibiting cellular senescence in a subject.

Claim 77 has been placed in both Groups II and III. The antibody embodiment will be searched with Group II. The nucleic acid embodiment will be searched with Group III.

Claim 79 has been placed in both Groups VIII and IX. The plant embodiment will be searched with Group VIII. The animal embodiment will be searched with Group IX.

Claims 83 and 85 have been placed in both Groups XI and XII. The plant embodiment will be searched with Group XI. The animal embodiment will be searched with Group XII.

The inventions listed as Groups I-XIV do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Group I is directed to purified lats protein, analogs, fragments, chimeric constructs, to the DNA that encode them and to a pharmaceutical composition and kit, which is the first appearing product, method of making and method of using. The special technical feature is the disclosed protein and DNA sequences. Group(s) II-III, XI-XII are drawn to structurally different products which do not share the same or a corresponding technical feature. Group(s) IV-X and XIII-XIV are drawn to methods having different goals, method steps and starting materials, which do not share the same or a corresponding special technical feature. Note that PCT Rule 13 does not provide for multiple products or methods within a single application. Since the special technical feature of the Group I invention is not present in the Group I claims, and the special technical features of the Group II-XIV inventions are not present in the Group I claims.



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